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NATIONAL INSTITUTE FOR OCCUPATIONAL SAFETY AND HEALTH

convenes

MEETING TWO

WORLD TRADE CENTER HEALTH PROGRAM
SCIENTIFIC/TECHNICAL ADVISORY COMMITTEE

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STEVEN RAY GREEN AND ASSOCIATES
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February 15, 2012

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TRANSCRIPT LEGEND

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In the following transcript: a dash (--) indicates an unintentional or purposeful interruption of a sentence. An ellipsis (. . .) indicates halting speech or an unfinished sentence in dialogue or omission(s) of word(s) when reading written material.

-- (sic) denotes an incorrect usage or pronunciation of a word which is transcribed in its original form as reported.

-- (phonetically) indicates a phonetic spelling of the word if no confirmation of the correct spelling is available.

-- "uh-huh" represents an affirmative response, and "uh-uh" represents a negative response.

-- "*" denotes a spelling based on phonetics, without reference available.

-- (inaudible)/ (unintelligible) signifies speaker failure, usually failure to use a microphone.

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P R O C E E D I N G S

(12:08 p.m.)

WELCOME

DR. MIDDENDORF: Good afternoon. If the committee members will come to the table, appreciate it, we'll get started. I have a few administrative details that we need to take care of here at the beginning. I'd like to extend a warm welcome to the members of the public who are here in the room and also those who are on the phone. We very much appreciate your interest in these proceedings and look forward to your participation. For those who have signed up who would like to make comments, we do have public comments scheduled to begin at 3:45 this afternoon and then we'll have another public comment session tomorrow morning.

For those of you who are here in the room, I'll point out the emergency exit routes. If you look around the room, you'll notice that there are three doors that have exit signs above them. You need to ignore two of those exit signs. The exit sign back here behind me to the left is not an exit door. Please don't

1 go out that way.

2 The double doors in the back far corner of
3 this room are not exit doors. Please do not
4 go out of those either. If, for some reason
5 we need to evacuate the room, this door that's
6 about three quarters of the way down here on
7 my left is the door to go out. And the
8 quickest way to get out is when you go through
9 that door, turn to your right, go until you
10 see two double glass doors on your left. Go
11 through those double glass doors, immediately
12 turn right, go down that hallway, and you'll
13 see a door that says Fire Exit on it, and
14 that's the way you get out of the building.
15 So please, that would be the best way to do
16 it.

17 For those of you on the phone, I suggest that
18 you look around, figure out the evacuation
19 route for your buildings. I need to point out
20 that we do have copies of the agenda for this
21 meeting. They are on the back table, and
22 they're also available on the committee's
23 website for anyone who is on the phone. You
24 can download the agenda from our website.
25 We also have copies of the public comments

1 that were received as of about 11 on February
2 13th. They have been offered, filed to the
3 committee before the meeting, and they're here
4 on the back table. If you don't want to haul
5 around a lot of paper with you, these comments
6 will be posted on NIOSH's docket, which is
7 docket number 248 for this committee and
8 that's also available through the committee's
9 website.

10 We need to do a quick roll call, and so we'll
11 go around the table first and I'd ask each of
12 the members to identify themselves and state
13 whether or not there have been any changes in
14 their employment or interest that would affect
15 their conflicts of interest, and then we'll go
16 to the members on the phone.

17 This is going to be a little difficult because
18 we only have two working microphones.

19 MS. MEJIA: Good afternoon. Guillermina
20 Mejia, no changes.

21 DR. QUINT: Julia Quint, no changes.

22 DR. ROM: Bill Rom, no changes.

23 MS. FLYNN: Kimberly Flynn, no changes.

24 MS. HUGHES: Catherine McVay Hughes, no
25 changes. I'll bring the mic over.

1 DR. TRASANDE: Leonardo Trasande, no changes.
2 DR. MARKOWITZ: Steven Markowitz, no changes.
3 MS. DABAS: Valerie Dabas, no changes.
4 MR. CASSIDY: Stephen Cassidy, no changes.
5 DR. NORTH: Carol North, no changes.
6 DR. TALASKA: Glenn Talaska, no changes.
7 DR. ALDRICH: Tom Aldrich, no changes.
8 DR. HARRISON: Bob Harrison, no changes.
9 DR. WARD: Liz Ward, no changes.
10 DR. MIDDENDORF: Okay, and -- oh, I'm sorry.
11 MS. SIDEL: I'm Susan Sidel, no changes.
12 DR. MIDDENDORF: Thank you, and on the phone?
13 DR. DEMENT: John Dement, no changes.
14 DR. WEAVER: And Virginia Weaver, no changes.
15 DR. MIDDENDORF: Okay, thank you all very
16 much. To those of the members who are on the
17 phone, please let me know when you leave and
18 when you return so we can be certain that we
19 continue to have a quorum.
20 Also, I want to remind everybody that there
21 may be some topics which come up that present
22 a conflict of interest for members. And when
23 these topics come up, I'll ask each of the
24 members to state that they are recusing
25 themselves so we have that on the record.

1 That's just the best way to handle that.

2 I also ask everybody to -- we have a couple of
3 issues; one is the microphones. We only have
4 two microphones available in this room.

5 Tomorrow we will be moving into conference
6 rooms A and B, so we'll have more microphones
7 in there. We're going to leave this
8 microphone turned on so we don't have that
9 problem with the lag time that we had before,
10 and then we'll just pass it around. I just
11 wanted to point that out.

12 One of the microphones will be up at the
13 podium until we're done with presentations, or
14 if presenters want to present from their
15 table, they can do that and we'll just give
16 them that one from the podium. I think that's
17 all I need to handle right now, so I will turn
18 this over to our chair, Dr. Ward.

19 DR. WARD: Good afternoon. The first speaker
20 today will be Dr. John Howard. He will give
21 us introductory remarks.

22 **INTRODUCTORY REMARKS**

23 DR. HOWARD: Can you hear me? Good afternoon.
24 Welcome to the second meeting of the
25 Scientific Technical Advisory Committee for

1 the World Trade Center Health Program. It is
2 with sadness that we begin our meeting.
3 Today, not only noting the passing of
4 responders and survivors since September 11th,
5 2001, but also the recent passing of Dr.
6 Stephen Levin, Professor of Preventive
7 Medicine at the Mount Sinai School of
8 Medicine.
9 For over 40 years, Dr. Levin treated,
10 counseled, and fought for thousands of
11 patients who were ill because of hazardous
12 exposures in their workplace. As Co-director
13 of the World Trade Center Worker and Volunteer
14 Medical Screening Program at Mount Sinai, he
15 was an early and prominent figure fighting for
16 a long-term health program to identify and
17 treat individuals who worked or volunteered at
18 the former World Trade Center site.
19 For all of his tireless work on behalf of the
20 World Trade Center Health Program during its
21 earliest and most difficult time, we honor him
22 and his service to his patients, to the City
23 of New York, his country, and to all of us.
24 Please join me in a moment of silence to honor
25 the recent passing of responders, survivors,

1 and Dr. Levin.

2 (pause)

3 I have four items for you today before we
4 begin the meeting. The first item is the
5 teleconference meeting on January 24th. I
6 apologize for the technical problems which
7 caused the cancellation of the 24th January
8 teleconference meeting of the committee. We
9 are taking steps to ensure there will be no
10 repeat of the technical problems if the
11 committee should want to hold another
12 teleconference meeting in the future.
13 Second, during this meeting, you will hear a
14 report regarding scientific findings and
15 support for establishing the statutorily
16 required criteria for Pentagon and Shanksville
17 responders. Commander Robert McCleery of the
18 NIOSH Division of Surveillance, Hazard
19 Evaluations and Field Studies in Cincinnati,
20 Ohio has provided a report which you have
21 already received and today will make a
22 presentation regarding his research on the
23 potential eligibility criteria for these
24 groups of responders.
25 I want to thank you in advance for your

1 consultation on this important issue. Please
2 note that no formal written communication from
3 the committee on eligibility criteria is
4 required. The meeting transcript will
5 suffice.

6 Third, I also appreciate the committee's
7 continuing consultative thoughts on research
8 needs for the World Trade Center Health
9 Program. Your thoughts to date have been
10 extremely helpful. And in addition to the
11 formal research funding announcement from the
12 NIOSH Office of Extramural Programs, the
13 committee's views about important knowledge
14 gaps and research needs will be placed on the
15 World Trade Center Health Program's website
16 for potential researchers to review.

17 Again, thank you in advance for your
18 consultation on this important issue. Please
19 also note that no formal written communication
20 from the committee on research needs is
21 required. The meeting transcript will
22 suffice.

23 Fourth, as you continue your discussion of
24 Petition 001 to add cancer or types of cancer
25 to the list of World Trade Center-related

1 health conditions, please keep in mind that
2 the Zadroga Act in Section 3312(a)(6)(C) notes
3 that the advisory committee must submit their
4 recommendation on the petition to the
5 administrator within 60 days or by a date
6 specified by the administrator, not to exceed
7 180 days from the date of the administrator's
8 request.

9 A request for a recommendation on Petition 001
10 was made to the committee on October 5th, 2011.
11 The maximum 180-day period for the committee's
12 consideration of Petition 001 ends on April
13 2nd, 2012. I had asked the committee to
14 provide its recommendation by March 2nd, 2012,
15 in order to provide enough time for the
16 committee chair to prepare the committee's
17 advice to the administrator.

18 However, since the opportunity for the
19 committee to meet on January 24th, 2012, was
20 cancelled, I would consider modifying the due
21 date for the committee's recommendation. If
22 the committee believes that more time is
23 necessary to reach a recommendation, I would
24 ask you to discuss that issue at this date and
25 for the chair to send a written request to me

1 for more time by the close of this meeting on
2 16 February.

3 Any additional discussion on Petition 001
4 after 16 February, 2012, must occur in another
5 public meeting, so please keep in mind
6 scheduling issues when determining whether
7 additional time would be beneficial to the
8 committee. In any case, the April 2nd due
9 date for a recommendation is a statutory
10 requirement; and therefore, no extension
11 beyond April 2nd can be approved.

12 I thank you again for your service. I wish
13 you a successful meeting.

14 **RESEARCH NEEDS**

15 DR. WARD: Okay. So, Rob McCleery has not
16 dialed into the call yet, so we're going to go
17 on and discuss research needs and then go to
18 Rob when he dials in.

19 So, I hadn't really planned a lot of
20 discussion around the research needs since I
21 think you've all seen the letter that we
22 prepared. But I didn't know if there were any
23 topics that any of you wanted to discuss
24 regarding the research needs or the conflict
25 of interest.

1 Oh, sorry, he's just gotten on the line, so
2 we'll proceed as planned with Rob McCleery's
3 publication -- I mean presentation.

4 **PENTAGON AND SHANKSVILLE, PA ELIGIBILITY**

5 MR. MCCLEERY: I apologize for that. I didn't
6 have this particular number, so I, again, I
7 apologize. So, good afternoon everyone.
8 Again, my name is Robert McCleery. I'm an
9 industrial hygienist at NIOSH here in
10 Cincinnati, Ohio. I appreciate the
11 opportunity to speak with you this afternoon
12 concerning the Pentagon and Shanksville,
13 Pennsylvania responses to the terrorist-
14 related aircraft crashes of September 11th,
15 2001.

16 Next slide, please. As it pertains to the
17 Pentagon and Shanksville sites, the World
18 Trade Center Health Program administrator is
19 required, conditioned to other
20 responsibilities to 1) determine the end dates
21 of cleanup at both sites and 2) determine
22 eligibility criteria relating to an increased
23 risk of developing a World Trade Center-
24 related health condition resulting from
25 exposure to airborne toxins, other hazards, or

1 adverse conditions resulting from the 9/11
2 terrorist attacks.

3 In the following slides, I will provide
4 information that addresses both of these
5 required determinations for the four
6 responding groups listed in the Zadroga Act
7 for the Pentagon and Shanksville sites: fire
8 department employees, police department
9 employees, recovery or cleanup workers and
10 contractors, as well as volunteers.

11 Next slide. At the Pentagon, fire department
12 personnel arrived on scene very shortly after
13 the aircraft crashed. Personnel within the
14 Arlington County Fire Department served as the
15 incident commanders during the fire rescue
16 phase of the response.

17 Numerous other fire departments responded to
18 the incident by backfilling other fire
19 stations or responding directly to the
20 Pentagon. This was set into action by mutual
21 aid agreements previously established between
22 these fire departments.

23 On September 21st, Arlington County Fire
24 Department transferred control of the site to
25 the FBI. The site now entered into the crime

1 scene phase of the response. At this time,
2 one firefighter company, a technical rescue
3 team, and paramedics remained at the site
4 until the FBI turned it over to the Department
5 of the Defense on September 26th or 28th.

6 The literature differs as to the date of
7 transfer of this command. From September 26th
8 or the 28th, the available literature does not
9 provide any information as to what period of
10 time fire department personnel were on site
11 until the end of the demolition and cleanup
12 phase of the incident on November 19th, 2001.

13 Next slide. The police departments. The lead
14 law enforcement agencies on site included the
15 Arlington County Fire Department, with
16 jurisdiction of areas surrounding the
17 Pentagon, Defense Protective Services, federal
18 law enforcement agencies within the Pentagon,
19 with jurisdiction of the Pentagon, and the
20 FBI.

21 Many other police departments respond --
22 responded either at the Pentagon or by
23 backfilling police stations, by way of the
24 Northern Virginia Law Enforcement Mutual Aid
25 Agreement or the Northern Virginia Sheriffs

1 Mutual Aid Agreement.

2 The available literature indicates that the
3 Pentagon response had a police department
4 presence until the FBI turned the site over to
5 DOD on September 26th or 28th, 2001. The
6 literature suggests that while the Pentagon
7 site was under DOD control, services typically
8 provided by police departments were handled by
9 military police or Defense Protective Service
10 personnel.

11 However, the literature does not provide
12 additional information as to what period of
13 time police department personnel were on site
14 until the end of the demolition cleanup phase
15 of the incident on November 19th, 2001.

16 Next slide. The Pentagon response and initial
17 cleanup of areas of the Pentagon surrounding
18 the incident site as employees began returning
19 to work on September 12th, 2001. The
20 demolition cleanup of the incident site itself
21 was delayed until after a memorial service
22 recognizing the one-month anniversary of the
23 9/11 attack on October 11th, 2001.

24 The demolition and cleanup activity of the
25 most severely impacted area began on October

1 18th, 2001, and concluded on November 19th,
2 2001.

3 Next slide, the volunteers. The information
4 in the literature does not provide a
5 comprehensive list of all of the volunteers
6 onsite for the time frames of participation of
7 those that did respond. Literature indicates
8 that there were many volunteers that played a
9 role in the response, with specific mention of
10 the Red Cross and Salvation Army.

11 It is reasonable to conclude at least some
12 volunteers were onsite through the FBI
13 relinquishing the site to DOD on September
14 26th or 28th, 2001. The literature does not
15 provide additional information pertaining to
16 volunteers remaining onsite through the
17 demolition and cleanup phase of the response.

18 Next slide. So the available information
19 concerning the Pentagon response does have
20 limitation. The information has uncertainties
21 as to when each of the responding groups faced
22 increased-risk activity at the Pentagon site.

23 Next slide. For the Pentagon response to the
24 September 11th terrorist-related aircraft
25 crash, the recommended concluding date is

1 November 19th, 2001. To ensure that each of
2 the groups that did respond are provided
3 adequate opportunity for medical monitoring
4 and treatment benefits, the World Trade Center
5 Health Program eligibility is recommended for
6 the period covering September 11th, 2001
7 through November 19th, 2001.

8 The available literature indicates that
9 documented air and wipe sample monitoring
10 conducted through September 28th, 2001, did
11 not reveal any exposures of concern. However,
12 no information is available on exposures
13 during the demolition of areas directly
14 affected by the aircraft crash.

15 The next few slides will cover the
16 Shanksville, Pennsylvania response. Next
17 slide, please. At the Shanksville site, fire
18 department personnel arrived onsite shortly
19 after the aircraft crashed. The FBI
20 controlled the site from the onset of the
21 response. Most of the fire department
22 personnel left the site after the FBI turned
23 the site over to the Somerset County coroner
24 on September 24th, 2001.

25 There was a limited fire department presence

1 until the conclusion of the final sweep of the
2 crash site which took place on September 29th
3 and 30th, 2001. The available information
4 does not indicate whether fire department
5 personnel were onsite during the site
6 restoration activity from October 1st through
7 October 3rd of 2001.

8 Next slide, Shanksville Police Department.
9 Law enforcement personnel were also on site
10 quickly after the aircraft crashed. Like the
11 fire department, most police department
12 personnel left the site after the FBI
13 relinquished the site to the county coroner.
14 Police department presence was limited at the
15 Shanksville site until the conclusion of the
16 final sweep of the crash site for aircraft
17 parts and potential human remains on September
18 29th and 30th, 2001.

19 The available information does not indicate
20 whether police department personnel were on
21 site during the site restoration activities
22 from October 1st through the 3rd of 2001. The
23 literature does suggest that law enforcement
24 personnel remained at the Shanksville site for
25 a number of years to provide security.

1 Next slide. For the recovery or cleanup
2 contractors, the literature indicates that
3 environmental restoration contractors restored
4 the site as close as possible to the original
5 appearance as they could from October 1st
6 through the 3rd, 2001.

7 This included backhoeing the crater with soil,
8 adding topsoil to the crater area as well as
9 the forested area near the site and seeding
10 the area with flowers and grasses.

11 Next slide, volunteers. The available
12 information does not provide a comprehensive
13 list of all of the volunteers onsite or the
14 time frames of participation of those that did
15 respond. The Red Cross and Salvation (sic)
16 are cited as responding to the Shanksville
17 site. Like fire and police personnel, most of
18 these volunteers left the site on September
19 24th, 2001 and had limited presence until the
20 final sweep of the site on September 29th and
21 30th.

22 The available information does not indicate
23 whether volunteers were on site during the
24 October 1st through the 3rd site restoration
25 activity. As with the Pentagon, the

1 Shanksville site has limitations in the
2 information and that information has
3 uncertainties as to when each of the
4 responding groups ceased increased risk
5 activity at the Shanksville site.

6 Next slide. The Shanksville response to the
7 September 11th terrorist-related aircraft
8 crash, the recommended concluding date is
9 October 3rd, 2001. And to ensure that those
10 who did respond were provided adequate
11 opportunity for medical monitoring and
12 treatment benefits, the World Trade Center
13 Health Program eligibility recommended for the
14 period covering September 11th, 2001 through
15 October 3rd, 2001.

16 Environmental monitoring at the site indicated
17 that surface soil, subsurface soil, and
18 groundwater did not exceed Pennsylvania
19 Department of Environmental Protection health
20 standards. Remediation was not required at
21 the site. No indication that surface water
22 contamination was attributable to the crash.

23 Next slide. The following is proposed
24 eligibility criteria for the Pentagon
25 responder: being a member of the fire or

1 police department, whether fire or emergency,
2 active or retired or worked for a recovery or
3 cleanup contractor or was a volunteer who
4 performed rescue, recovery, demolition, debris
5 cleanup, or other related services at the
6 Pentagon site for terrorist-related aircraft
7 crashes of September 11th, 2001 for at least
8 one day during the period beginning September
9 11th, 2001, ending on November 19th, 2001.

10 Next slide. The following is the proposed
11 eligibility criteria for the Shanksville
12 responder: member of a fire or police
13 department whether fire or emergency, active
14 or retired or worked for a recovery or cleanup
15 contractor or was a volunteer who performed
16 rescue, recovery, demolition, debris cleanup
17 or other related services at the Shanksville,
18 Pennsylvania site for the terrorist-related
19 aircraft crash of September 11th, 2001, for at
20 least one day during the period beginning
21 September 11th, 2001, and ending on October
22 3rd, 2001.

23 This concludes my presentation for this
24 afternoon.

25 DR. WARD: Are there questions for Rob? So,

1 does anyone on the committee want to ask any
2 questions or make any comments about Rob's
3 presentation?

4 DR. HARRISON: Thank you very much for all the
5 comments. I think it's very reasonable.

6 DR. WARD: I agree. Is that the general sense
7 of the committee, that it's reasonable? Okay,
8 well, we'll record that for the record.

9 **RESEARCH NEEDS**

10 So, now we'll go back to the research needs
11 and where we were on that was I was asking if
12 anyone had any questions or felt the need for
13 more discussion on the research
14 recommendations or the document that was
15 circulated regarding principles for handling
16 conflict of interest within this committee.

17 **PETITION ON CANCER**

18 Okay, hearing none, we'll move on, and I guess
19 our next topic is the petition on cancer. For
20 those on the phone, I am going to be moving to
21 the podium so that I can present some slides I
22 prepared, and that will take -- that
23 transition will take just a minute. It will
24 be another minute because Paul is conferring
25 on something. Are we okay to proceed?

1 Well, I think as most of the committee members
2 know but possibly some members of the public
3 may not, we had hoped to discuss -- is this
4 on? Is that better?

5 DR. MIDDENDORF: Would you prefer to use this
6 one or that one?

7 DR. WARD: Maybe we should use the other one,
8 and probably we should turn this one off.

9 Thank you. I do have a small voice, so this
10 will be very helpful.

11 As most of you know, when we had to -- when we
12 weren't able to have our last meeting by
13 teleconference, one -- the plans for how we
14 were going to address the petition on cancer
15 was one of the things that we were going to
16 discuss as a committee, so in the absence of
17 having that meeting, I really thought hard
18 about how we could approach this topic in a
19 way that we could really have meaningful
20 discussion at this meeting despite that
21 circumstance.

22 And as you all know, we received a letter from
23 Dr. Howard subsequent to a letter he received
24 from several congressmen asking us to review
25 the available information on cancer outcomes

1 associated with exposure resulting from the
2 September 11th terrorist attacks and provide
3 advice on whether to add cancer or a certain
4 type of cancer to the list of World Trade
5 Center-related conditions.

6 And as we discussed that at our last meeting,
7 I think we realized that there were a number
8 of very complex and difficult questions
9 embedded in that -- in that request. And one
10 of them was basically whether, based on what
11 people were exposed to at the World Trade
12 Center, do we believe it's possible, probable,
13 or not that the exposures could cause cancer.
14 And it's -- whatever our recommendation is, we
15 would need to provide a scientific rationale.
16 Now there's a second topic. There's at least
17 one other really complex topic that came up at
18 our last meeting, which was what are the
19 criteria for having a health condition?

20 And so my idea was to focus today's
21 presentations and discussion on the first
22 question: Do we believe it's unlikely,
23 possible, probable, et cetera, that exposure
24 to the dust may cause cancer, and then
25 depending on where the committee stands at the

1 end of the day, we'll decide how best to use
2 our time tomorrow.

3 And I think it's important. My boss says --
4 at the American Cancer Society -- says this
5 all of the time, so I guess he's implanted it
6 in my head. I think when we talk about the
7 scientific rationale, it's really going to be
8 important to talk about what we know, what we
9 don't know, and what we believe, because I
10 think that, you know, we'll all -- in all the
11 presentations today, one recurring theme will
12 be we wish we had more data; we wish we
13 understood the exposures better; we wish we
14 knew more.

15 **EPIDEMIOLOGY AND OVERVIEW OF MECHANISMS OF CARCINOGENESIS**

16 So what I'll be doing is just reviewing the
17 epidemiologic studies that are completed and
18 ongoing. I am going to talk about the
19 potential carcinogens present in the World
20 Trade Center dust, and then I am going to give
21 a quick overview on mechanisms of
22 carcinogenesis, really focusing on those
23 issues that I think pertain most to our
24 discussion today.

25 So with respect to the epidemiologic cohorts,

1 we had several presentations on them at our
2 last meeting and we also have access to
3 published information on them. So I am just
4 going to go through them very quickly.

5 Among the cohorts that are under study, there
6 are -- there's studies going on of the Fire
7 Department of New York, and I think these
8 studies probably from an epidemiologic point
9 of view are the most -- are going to be the
10 most complete and informative because we know
11 that they really have a well-defined
12 population and a population that is, you know,
13 highly exposed, a comparison group.

14 And they also have a separate set of EMS
15 workers that has not been published on.

16 They're also doing an employer-based medical
17 screening program, which will provide
18 additional information.

19 The second large cohort that can be studied is
20 the New York -- is from the New York and New
21 Jersey World Trade Center Clinical Consortium,
22 and I think that will also be a very
23 informative study. It will suffer from the
24 limitation that it essentially was a self-
25 referred group of people.

1 The third one, which I'm not sure is actually
2 being studied for cancer or not. I'm sure
3 someone in the room knows. It's the cohort
4 that's been identified through the World Trade
5 Center Environmental Health Center, and this
6 population is unique because it includes some
7 children.

8 And then there's the very large World Trade
9 Center Health Registry that's being run by the
10 New York Health Department. And that one is
11 clearly the largest in terms of sample size.
12 Probably the most severe limitation is that
13 about 70 percent of the cohort is self-
14 referred rather than identified from the list
15 or records, and that group is being followed
16 both by surveys and by linkage with cancer
17 registries and mortality data.

18 So in the first publication of cancer
19 incidence data from the firefighters cohort,
20 the incidence ratio for all cancers combined
21 was 1.10 compared to the general population.
22 And depending on particular adjustments used,
23 it was 1.19 to 1.32 in comparison to non-
24 exposed firefighters.

25 There are also some excesses for particular

1 cancer sites. The findings differed a little
2 bit based on which adjustment was used, but
3 basically, there were significantly elevated
4 or borderline excesses observed for stomach,
5 colon, melanoma, prostate, thyroid, and non-
6 Hodgkin lymphoma compared to the general
7 population rates.

8 And I think one thing that's important to note
9 here, because it's been noted by others in the
10 literature, is that there are a number of
11 these cancers that no -- are likely to be
12 detected by screening or by just access to
13 medical care, and the paper did attempt to
14 control for that bias in the analysis.

15 But with respect to other epidemiologic
16 studies, in the first publication from the
17 World Trade Center Health Registry study,
18 there was no excess of all cancers combined or
19 eight major organ systems reported. There
20 have also been case reports suggesting the
21 possible excess of multiple myeloma in the
22 literature.

23 So I think one of the things that it's
24 important to understand before we move on from
25 the epidemiology studies is that epidemiologic

1 studies in general have their strengths and
2 their weaknesses. One of the strengths is
3 that you're actually studying the events, not
4 animal systems or models.

5 On the other hand, it's often very difficult
6 in epidemiologic studies to accurately
7 estimate exposure, and I think that applies
8 even more so in these studies; although, I
9 think there have been really good attempts to
10 use surrogates of exposure, like in the
11 firefighter cohort, kind of developing
12 exposure classifications based on when people
13 arrived at the site, for example.

14 So I think that the existing studies are doing
15 the best job that they can, but ideally, you
16 know, what you'd love is an exposure matrix
17 for each person so that you knew, you know,
18 this person was very highly exposed and they
19 didn't work well. And that's probably not
20 going to be present.

21 And so, when you don't have good exposure
22 information, you may not be able to see some
23 of the things that you tend to look for when
24 we look for causal association, so we may not
25 see a strong dose response, because we don't

1 have good exposure data. We may not see the
2 trends that one might expect to see.
3 Another criteria for causality that's
4 considered is consistency between studies, and
5 again, I think, especially in this case, we
6 may not see that level of consistency because
7 we don't have one exposure. We have many
8 exposures, and we have different populations
9 and individuals who were exposed to different
10 things, so I would not be surprised at all
11 with the different studies that they show
12 increased risk for cancer. They may see
13 increases at different sites, so I think we
14 have to be really cautious about especially
15 making negative conclusions about the findings
16 of these studies.
17 And the last -- well, the last one on this
18 slide is even though many of these populations
19 are sizable, they're still, in many cases,
20 small enough or early enough in the follow-up
21 period that there are not very many cases
22 expected based on population rates.
23 So if we don't see an effect, we really need
24 to be careful in interpreting that because it
25 may be -- the studies may be too small to rule

1 out small risks or risks for rarer cancers.
2 One of the most important things, and I know
3 it came up in our discussions last time, and
4 I'm sure it will come up again today, is that,
5 you know, I think when we all were trained in
6 occupational health, those of us who were, we
7 all thought, well, you know, usually solid
8 tumors you're looking for at least 20 years
9 between the onset of exposure and disease and
10 hematologic cancers, the latency period is
11 shorter.

12 And -- but I guess what I wanted to emphasize
13 is the issue of latency period is most
14 relevant in epidemiologic studies early in the
15 follow-up period when we have negative results
16 and follow-up may be too short to see a
17 positive effect.

18 It's not necessarily relevant in the sense of
19 saying, well, this cancer can't be related to
20 exposure because, you know, the exposure only
21 occurred five years ago. I'll get more into
22 that later, but I don't think you can make
23 those kinds of assumptions based on what I'll
24 present to you about the mechanisms of
25 carcinogenesis.

1 So, if -- I think we got the -- I got the
2 sense in the discussion last time, and this
3 doesn't probably represent everyone's
4 viewpoint, but I did get the sense from the
5 discussion that many people felt that they
6 could not make a decision on the cancer
7 petition based on the epidemiologic data
8 alone.

9 Obviously, the strongest study is the
10 firefighter study, but I don't -- I didn't
11 sense an overwhelming consensus that the
12 findings of that study were so definitive that
13 it would be the basis for a recommendation.
14 So then the question was, what can we learn
15 from looking at the exposure data, but I think
16 we have to acknowledge at the outset that it's
17 incredibly difficult to interpret the --
18 especially air sampling data from the World
19 Trade Center study.

20 And one critical limitation was that there's
21 almost no data from the first week after the
22 attack. A lot of people said that last time,
23 and I think, you know, I think we all
24 understand that. I'm puzzled about some of
25 the air data, because it really seems like the

1 low air levels measured in some of the
2 personal air sampling studies done on the
3 workers seems really inconsistent with the
4 extent of respiratory symptoms that we're
5 seeing.

6 And so I don't know how to answer that
7 question, but it's my belief that it's, you
8 know, I don't see it fitting together well.

9 So, one approach to looking at the cancer
10 hazard which I thought we could take today is
11 really to focus on the composition of the
12 initial dust and smoke as reflected in the
13 mass dust samples that were collected.

14 And those samples were collected and analyzed
15 by more than one group so at least we have
16 some -- we can look at consistency of their
17 findings. And the other benefit, I think, of
18 looking at the dust and smoke is that there
19 were a lot of populations exposed to that.

20 So, for example, we know that there were fires
21 at the site, and we knew that -- we know that
22 firefighters and police officers who were on
23 the site itself were exposed to combustion
24 products from the fires, but just for the
25 purposes of having a simpler discussion today

1 and a discussion that kind of encompasses
2 exposures to all of the groups, I thought we
3 could first focus on the dust and smoke,
4 recognizing that there's more -- there's more
5 to the story that we'll have to get to later.
6 So, in poring through the literature and, you
7 know, all of the exposure papers, I have to
8 confess, I am not a chemist; I am not an
9 industrial hygienist, and it's not easy to
10 read these papers. But, you know, one of the
11 things that I got out of it was really, you
12 know, what went into the buildings is really
13 what came out of the buildings.

14 So, if you look at, you know, there was a lot
15 of light-weight concrete; there was asbestos;
16 there was gypsum; there was drywall; lots of
17 glass. There was glass fragments and man-made
18 vitreous fibers from insulation. We know that
19 there were polycyclic aromatic hydrocarbons
20 measured in the bulk samples. We know that
21 there were metals measured in the bulk
22 samples.

23 And then, we also know that there were
24 volatile organic compounds in the mix. Now
25 those probably, looking at the dust, is not

1 the best way to look at exposures to those,
2 which is why I have them in blue, because we
3 know they were there. In the dust, though,
4 they may have been absorbed onto particles and
5 fibers and other things, so they may be there,
6 but it's probably not the best way to measure
7 them.

8 So, what, I mean, what -- so, two of the
9 reasons I focused on these particular
10 exposures is one, that they were pretty
11 substantial. So, for example, the asbestos
12 was, you know, in a few of the bulk samples
13 was from .8 to 3 percent of the total weight
14 of the sample. So that's pretty significant.
15 The other thing is a number of them are --
16 have been recognized as human carcinogens for
17 which, based on epidemiologic data, so they
18 are substances for which we have fairly strong
19 epidemiologic data.

20 So that's why we're focusing on these
21 particular exposures. It doesn't mean that
22 there aren't other classes of exposures of
23 concern, and you know, we're not talking today
24 too much, at least in the presentations, about
25 PCBs and furans and, you know, TCDDs, but

1 again, you know, we have a limited amount of
2 time, and I wanted to focus on the things
3 where I thought there was the clearest data to
4 talk about.

5 So, now shifting gears a little bit, and I
6 want to thank both Julia and the National
7 Cancer Institute for these slides. Julia
8 pointed out to me that there was a slide set
9 on the National Cancer Institute website that
10 we could use for this presentation because I
11 think that a picture is worth a thousand
12 words.

13 So all of the slides in blue come directly
14 from that website and have not been modified.
15 So basically, what is cancer? So, when a cell
16 becomes cancerous, basically, it loses the
17 ability to control its own growth and to
18 organize itself appropriately in tissues. And
19 this -- one of the key things in that process
20 is the damage to the DNA of the cell.

21 So this is a slide that summarizes a number of
22 different characteristics of cancer cells, and
23 it's really, at least historically the way
24 that cancer has been recognized is
25 pathologists look under a microscope at the

1 appearance of the cells from the tumor. So
2 the cells will be different. They'll have
3 larger nuclei. They will not organize
4 themselves into neat structures the way
5 they're supposed to.

6 So that's a real quick review of that, but
7 you, typically, you know, for our classic
8 carcinogens, both tobacco and asbestos, we see
9 a 20-year latency period, and that's -- but
10 what that means is in 20 years from the onset
11 of exposure to the peak of disease in the
12 population, so in this case, men started
13 smoking in the United States soon after 1900,
14 and we saw the peak in lung cancer in the
15 1970's.

16 So the -- so as I mentioned, the key, you
17 know, the critical step in carcinogenesis is
18 an interaction of exogenous or an endogenous
19 substance with DNA within the cell, and that
20 can be a chemical, it can be a virus, it can
21 be radiation. So there is a component where
22 there is an interaction with DNA.

23 And typically, what happens, and this is
24 grossly oversimplified, but basically the DNA
25 is the cell's mechanism that basically codes

1 for the production of everything a cell needs
2 to grow and sustain life. So, what happens is
3 when there's a chemical damage, for example,
4 that might change one of the -- and so, and
5 the code is really in the three -- it's in
6 three, you know, it's in three chunks.
7 So, CAA codes for a particular thing, and if
8 you substitute one of its -- one of the
9 chemicals, it changes the whole, that whole
10 code. So, basically, three things can happen.
11 You can change a single base. Those things
12 are called bases, and the three together are
13 (indiscernible).
14 You can change a base. You can put an
15 addition in a base, or you can make a deletion
16 from the base, but in any case, it basically
17 messes up the code such that the gene is not
18 effectively doing what it's supposed to do.
19 And there's really three kinds of genes that
20 are involved in the process of carcinogenesis.
21 One type -- and you know, this is large
22 categories. One type is oncogenes, and what
23 oncogenes do is they -- when they're -- they
24 accelerate cell growth and division. Tumor
25 suppressor genes enable the cell to put a

1 brake on that kind of uncontrolled growth and
2 DNA repair genes allow the cell to repair
3 errors or mutations in the DNA itself.

4 So what happens, if you're exposed to a
5 carcinogen and you have a mutation and in any
6 of those three types of critical genes, if the
7 cell does not repair that mutation before it
8 divides, that mutation is going to be passed
9 on to the daughter cells.

10 So typically what we see in cancers is
11 multiple mutations, and it's kind of, it's
12 thought that these mutations occur over a
13 period of time, so possibly, you know, when
14 you're 25, you get a mutation in a tumor
15 suppressor gene, and if that is maintained,
16 then as those cells divide and proliferate,
17 they accumulate additional mutations, and in
18 that process, though, you're not just -- the
19 changes in, the mutations in the genes is not
20 the only thing going on to lead to cancer.
21 Other things are going on that kind of promote
22 the growth of those cells.

23 So for example, for breast cancer, estrogen
24 promotes the growth of tumors in the breast
25 because breast tissues are naturally sensitive

1 to, you know, hormones, for example. So it's
2 not only the genetic mutation or the
3 interaction with the DNA. It's multiple
4 things going on.

5 And so, we tend to divide the process of the
6 carcinogenesis into four big buckets:
7 initiation, which is basically, at least an
8 initial mutagenic effect; promotion, which is,
9 you know, encouraging those abnormal cells to
10 grow; malignant transformation, which means
11 that the cell has kind of passed beyond the
12 point where it can revert back to a normal
13 cell. It's accumulated enough damage that
14 it's essentially destined never to go back to
15 normal. And then ultimately that tumor gets
16 larger and invades other tissues beyond where
17 it arose and it can metastasize to other parts
18 of the body.

19 So the reason I'm emphasizing the promotion
20 and progression is, is that it's important in
21 the context of the exposures we're discussing
22 today because inflammation is one of many --
23 it's one of the important mechanisms of
24 carcinogenesis. And inflammation actually can
25 do a large number of different things, but

1 basically inflammation is a normal response to
2 tissue damage that can result from infection,
3 chemical irritation, and/or wounding.

4 However, when it becomes chronic and it
5 becomes chronic in a number of known diseases,
6 it can damage the body and lead to illness.

7 So, for example, we've all heard of Crohn's
8 disease, which is kind of an inflammatory
9 condition of the bowel, cirrhosis of the
10 liver, which is an inflammatory condition of
11 the liver. Many of the diseases, especially
12 the infectious diseases that result in
13 inflammation also result in cancer.

14 And inflammatory processes can also occur as a
15 result of chronic chemical and mechanical
16 inflammation, but it's important to know that
17 inflammation in general can really lead to
18 cancer in a multitude of ways. Its increasing
19 cell proliferation and turnover is actually
20 generating mutagenic substances from some of
21 the reactions that release oxygen and nitrogen
22 species, and it's also producing cytokines and
23 growth factors and other biologically active
24 chemicals that are influencing the
25 microenvironment around the area where the

1 potential tumor is developing.

2 With regard to mechanism, I guess the other
3 things I wanted to mention are that -- one of
4 the things we have to consider is that for
5 many of the people in the exposure group, the
6 duration of actual exposure is relatively
7 short, but I think it's important to note that
8 at least in some of the populations studied,
9 inhaled fibers and dust can remain in the body
10 for a very long time. And so, in fact, a
11 short-term environmental exposure can lead to
12 a long-term biological exposure, and we've
13 seen that in some of the bronchial lavage
14 studies.

15 The other thing is, you know, we've talked
16 about this average latent period for solid
17 tumors, but I think it's important to
18 recognize that it all depends on what stage in
19 the cancer process an exposure occurs. So,
20 for example, we see this curve in the
21 population when in relation to onset of
22 smoking in the population at large, you know,
23 and then the lung cancer epidemic following 20
24 or 30 years later.

25 But when a person stops smoking, their lung

1 cancer risk goes down dramatically within
2 three to five years. So, what, you know, one
3 thing that's probably happening there is that
4 essentially tobacco smoking contains
5 practically every carcinogen known to man, and
6 some of those substances actually are
7 promoting or, you know, causing the tumor to
8 progress, so they're both initiators and
9 promoters.

10 And so you see this much more rapid effect in
11 an individual that stops smoking than you
12 would expect from the long latency period for
13 the initiation, and we've seen something
14 similar recently in breast cancer and this is
15 really interesting.

16 So, in 2002, the Women's Health Initiative
17 published a study showing that use of
18 postmenopausal hormone therapy was associated
19 with an increased risk of breast cancer and
20 the surveillance epidemiologists noted in that
21 year's data that there had been a dramatic
22 drop in breast cancer incidence virtually the
23 same month that those studies came out.

24 And at the time, you know, everybody was
25 saying it can't be related to HRT, it's not

1 biologically plausible that something could
2 act that fast. Well, if, you know, there's
3 pretty good consensus now. I don't think
4 anyone disagrees that one of the major factors
5 or the major factor in that abrupt decline is
6 that, you know, on a population basis, a lot
7 of women stopped taking HRT, and HRT was
8 really promoting or causing tumors to progress
9 in the women.

10 And since that time we've actually seen a
11 flattening out of rates. It's not continuing
12 to go down, which further supports the
13 hypothesis that it was that one time decline
14 in HRT.

15 So, we'll be moving on. I have a few more
16 things I'd like to present, but then we'll be
17 moving on to the presentations that I asked
18 people to prepare regarding specific exposures
19 of concern. But before I wanted to go on, I
20 wanted to mention that I think there is an
21 opportunity to learn more about the potential
22 health effects of the World Trade Center dust
23 exposure that maybe we haven't explored as
24 fully as we could.

25 So, one of the things I noticed in looking

1 through the literature is that, you know,
2 there was a lot of concrete in the buildings
3 and concrete is a -- you know, two of the main
4 components of concrete are cement dust and
5 silica. Silica, as I mentioned, is an
6 accepted lung carcinogen and it's also
7 associated with autoimmune diseases and stage
8 III lung disease.

9 Pulverized concrete also contains a material
10 called Portlandite, which is highly caustic
11 and not shown in this slide, but I know many
12 people in the room are aware of it. People
13 who work with wet concrete often get skin
14 sensitization because of hexavalent chromium
15 in the cement mix. And many European
16 countries actually regulate the content of
17 hexavalent chromium in their cement, but the
18 United States does not.

19 So -- but it appears, and again, this is very
20 preliminary -- it appears that maybe the
21 hexavalent chromium content of concrete once
22 it's set would not be as high as the
23 mesolithic form. But again, that is something
24 of concern.

25 But in fact, there have been a number of

1 studies of cement dust exposure, many of them
2 done, interestingly, in developing countries,
3 but many of these studies, and again, some are
4 small, but actually a few are, you know, large
5 enough and well designed, at least on the
6 surface. And many of the studies, not all,
7 find increased respiratory symptoms among
8 people who work in the production of cement,
9 and they also demonstrate reduced lung
10 function among people with long-term exposure.
11 What I found most interesting is that there
12 was one study that actually found an increased
13 risk of GERD-type symptoms among people
14 exposed to cement dust. And by the way, all
15 of these studies are on the FTP site under the
16 folder that says cement.
17 Of even more concern is there have been some
18 cohort case controlled studies that have
19 suggested associations between cement-exposed
20 populations -- and that could be either in the
21 manufacture or in the construction industry --
22 in cancer of the lungs, stomach, colon, head
23 and neck, pharynx and larynx.
24 So cement dust that has not been reviewed by
25 IARC or NTP and the only kind of official

1 review I could find of it on it popped up on
2 the web, and it seems to have been done by the
3 Health and Safety Executive of the UK, but the
4 version of the document online is a little odd
5 because it does not have a publication date.
6 It has a number, but no date, but I think it
7 was -- it looks like it was published in 2006.
8 And basically, their synthesis of the cancer
9 literature at that time was that the epi data
10 was not convincing, but that they felt that
11 some of the associations that had been seen
12 were biologically plausible in large part due
13 to the known inflammatory responses associated
14 with exposure to cement dust.
15 So one of the ways I thought -- I mean, I
16 thought I had a pretty reasonable way to frame
17 the discussion today and get into depth on
18 some of the most important issues, but I think
19 tomorrow, the agenda is wide open, and one of
20 the things I thought that might help us frame
21 an agenda would be to -- at the end of the
22 presentations, we'll kind of poll the
23 committee and ask each person to check one of
24 these words and turn them in -- so, this is
25 not a vote, it's just a poll.

1 And then what we'll do is we'll summarize the
2 distribution of the results, just kind of
3 arranged by the exercise. So, we'll summarize
4 the distribution of the results and that will
5 help us know, do we have two really different
6 viewpoints? Are some people really on the
7 side of probable proof and are other people
8 way off on unlikely, possible, or do we have,
9 you know, a distribution centered at the
10 middle?

11 And then we can really see, you know, how can
12 we use our time tomorrow to, you know, to see
13 if the group has a consensus or not or to
14 figure out what issues are of most, we're most
15 uncertain about. And again, we are all
16 prepared to tabulate these result in such a
17 way that you --

18 MS. HUGHES: I have a quick question. On the
19 slides --

20 DR. MIDDENDORF: Wait a minute.

21 MS. HUGHES: Hi, I have a quick question. On
22 the last slide, it says is the blank that
23 exposure World Trade Center may cause cancer.
24 Can we also use slash smoke, because not all
25 of the exposure was dust --

1 DR. WARD: Yes.

2 MS. HUGHES: Because not all of the exposure
3 was dust.

4 DR. WARD: Yes.

5 MS. HUGHES: Because then it would be more
6 consistent with some of the other slides.

7 DR. WARD: Yes.

8 MS. HUGHES: Okay, great, thanks.

9 DR. WARD: We can make that -- yeah. So,
10 anyway, I think this will be helpful in
11 framing tomorrow's discussion and, you know,
12 and these are various options that we could
13 discuss tomorrow. There may be -- it may be
14 that people feel that there's critical
15 evidence that we didn't cover today that we
16 should go into in more depth tomorrow.
17 It may be that there are clearly opposing
18 positions that we should try to address
19 tomorrow. If we're -- if there's apparently a
20 high degree of consensus, then we can start
21 talking about the rationale for the position.
22 If we are leaning towards saying probable,
23 then we can discuss the issue of what sites do
24 we think are probable, and then hopefully have
25 whatever -- wherever we are, and certainly we

1 can discuss the possibility of needing to have
2 another conference call or meeting before we
3 can make our recommendation.

4 So, with that, along with my presentation, are
5 there any questions?

6 DR. MARKOWITZ: So just a couple of comments.

7 One is I don't really favor taking a poll
8 before we have the public comments. We have
9 the public comments at the end of today and
10 beginning of tomorrow morning, because that
11 would add to the discussion, influence our
12 thinking, so I would advocate doing a poll
13 after that.

14 I would also like to have, you know, do some
15 discussion before we do a poll because I want
16 to hear what people think. So if you want to
17 do a poll, we could do it. We could change
18 the time, though, until tomorrow after public
19 comments and after there's at least some
20 initial discussion. I assume the purpose of a
21 poll is to sharpen further discussion.

22 Another comment I have is about the choices of
23 unlikely, possible, biologically plausible,
24 probable, definite, and that is that actually
25 I think biologically plausible stands with

1 both possible and probable, and so I'm not
2 sure that these are exclusive categories. And
3 I understand that it's preliminary, a rough
4 way of getting a sense, and I wonder whether
5 one alternative approach would be to consider
6 reasonably anticipated as a substitute for one
7 of the categories.

8 DR. WARD: Maybe probable?

9 DR. MARKOWITZ: Well a --

10 DR. WARD: I guess, that's the thing, it
11 sounds like probable to me but, so I guess if
12 -- we can make any changes that you all want
13 to make. It did occur to me that maybe the
14 timing was wrong, but again, the timing was
15 kind of thinking about how can we tabulate
16 these results so that we could leave people
17 thinking about how we're going to use our time
18 tomorrow.

19 And some people may even want to, you know,
20 think about ideas that they'd like to present
21 or do literature searches tonight, or, you
22 know, people could prepare to argue the main
23 points overnight and so I did -- well, I did
24 bring enough paper ballots that we could have
25 more than one poll, so that's one option.

1 Valerie?

2 DR. MIDDENDORF: I think Catherine had a --

3 MS. HUGHES: Yeah, I had a quick question.

4 DR. MIDDENDORF: So, Catherine, then Tom, then
5 Valerie.

6 DR. WARD: I think I need to have my eyes
7 transplanted so --

8 MS. HUGHES: I know we're all -- we're looking
9 at actually what was in the dust and what was
10 in the fumes. Are we going to look at also
11 the impact of the temperature, because it
12 wasn't as though the temperature was the
13 temperature of the day, because it was just so
14 hot. It was like 1000 degrees -- if people
15 were close would have been impacted and how
16 the items could have changed due to the
17 temperature, too.

18 DR. WARD: Yeah, and I think, you know, that
19 would fall under the category of things where
20 there's something that where there are
21 critical issues that we haven't discussed. I
22 don't know if anyone is prepared to talk about
23 the temperature today or, you know, has really
24 looked into that issue, but if you feel that
25 that's an important issue, we can see if

1 there's anyone who wants to comment on that
2 further or we can put it on a list of things.
3 Again, I guess the question is do we feel like
4 we have enough information to make a
5 recommendation now, or are there things that
6 we feel are so important that we need to wait
7 until, you know, somebody really studies them
8 well enough to talk about them.

9 I mean, I certainly couldn't talk about that
10 today, and I don't know if anyone else could.

11 DR. ALDRICH: I was going to suggest, if
12 there's going to be a poll, maybe two
13 questions: biologically plausible, yes or no;
14 and then the other four, pick one.

15 DR. WARD: Good.

16 DR. MIDDENDORF: We forgot Valerie.

17 MS. DABAS: Just because I am not a scientist,
18 I just want to get the definition of
19 biological plausibility just because I've seen
20 so many different ones on the websites.

21 DR. WARD: That's a good question. My
22 definition of it is that when you look at the
23 exposure and what was -- when you look at the
24 dust and smoke and you look at what was in the
25 dust and smoke, and you look at what the

1 toxicity of the, of that we've already
2 observed in the events and, you know, when you
3 look at all of those elements of data, it
4 makes sense that this exposure could cause
5 cancer based on what we know about the cancer
6 process and the components in the material.
7 Now, that's my definition. Someone else may
8 have a better one. Julia?

9 DR. QUINT: I think I agree with most of what
10 you said except I'm not limiting it to humans,
11 because I -- the animal data that shows that
12 something is carcinogenic, to me, means I
13 don't think -- there are only a few cancers in
14 animals that are not biologically plausible in
15 humans, so I think the animal data is a
16 plausible mechanism in humans as well.

17 DR. WARD: Yes, and I totally agree with that,
18 and --

19 DR. QUINT: I thought you did.

20 DR. WARD: Yeah. I am going to return to my
21 seat until we are done with --

22 MR. CASSIDY: Thank you. You've discussed a
23 lot of topics, and one that I think is
24 interesting when you look at this is, you
25 know, is it blank that the exposure to World

1 Trade Center dust may cause cancer, and I
2 think it's hard to, you know, may be hard for
3 some people to answer that unless you're
4 talking about a level of exposure, right?
5 So you were talking about cigarette smoke, and
6 I would think that the studies show if you
7 smoked one cigarette and stopped before you
8 had an exposure to tobacco that the likelihood
9 of developing something from that would be
10 different if somebody smoked five packs a day
11 for ten years, right?
12 So I think it's important that the part or at
13 least part of the discussion to the level of
14 exposure, and I tie that in to when you said
15 that the air sample data seemed to be
16 inconsistent. Well, the question is where was
17 that air sample data taken? And, you know, my
18 personal recollection is I didn't see anybody
19 standing on the Pile taking it.
20 So, I don't know where -- if they took it five
21 blocks away or ten blocks away or where they
22 took it. And on that note, the air sample
23 data, I would remind everyone that is -- there
24 was much discussion about whether or not that
25 was a political decision to say quote,

1 unquote, the air was safe because they wanted
2 to open up Wall Street. You know, we had to
3 get back to business, the country was shut
4 down. So, I just wanted to raise that point.
5 I think people that were there working at the
6 site knew the air wasn't safe no matter what
7 Christie Todd witnessed, so.

8 DR. WARD: Yeah, and I do want to, I mean, I
9 fully acknowledge those issues and I didn't
10 want to spend a lot of time on them today just
11 because I really feel like, you know, both the
12 committee discussions and the published
13 literature both, you know, essentially give
14 that same information. But it's really trying
15 to come up with other approaches that maybe
16 can be a little bit more revealing and make --
17 help us make a decision.

18 But I think, you know, there's at least,
19 there's a couple of exposure scenarios and I
20 think we should acknowledge that too so we
21 have people who were -- we have a very heavily
22 exposed group that was working directly on the
23 Pile, especially in the early time period. We
24 also have the potential for the community
25 residents and the workers to have prolonged

1 exposure to the dust that entered the homes
2 and office buildings.

3 Now, again, I don't know that you would expect
4 to see exactly the same health effects in
5 those two populations, but they're both
6 populations that may have significant
7 exposure, possibly to different substances and
8 different concentrations.

9 DR. MIDDENDORF: It's easy to forget that we
10 have some committee members who are on the
11 phone, out of sight, out of mind, so I just
12 want to ask if any members on the phone have
13 any questions or comments.

14 DR. WEAVER: I don't, but we're moving along
15 fairly quickly and I just want to point out
16 that I'll be teaching from 1:30 until 2:50 and
17 I'm scheduled to talk at 3:10, so, you know,
18 we can just juggle when I talk around class,
19 but when I am in class I'll have my cell
20 phone, so I can listen in.

21 MS. SIDEL: I just want to say that because we
22 don't have air samples from, you know, from
23 the day 9/11, that's why Officer Harris's
24 uniform is so fascinating, because it's like a
25 snapshot in time of what, what was there, and

1 I believe that this also -- another study of
2 what FDNY, I think, equipment that I've seen
3 that are also from the actual day 9/11 from
4 people that were working. So, you know, I
5 feel as though there's a lot of different air
6 samples and they sort of collectively say the
7 same thing, and that is that there were a lot
8 of carcinogens down there.

9 And then we start talking about, you know,
10 different zones of exposure, but you're never
11 going to get -- that's never also going to be
12 firm and there's definitely people that were
13 super-exposed, but then there's also other
14 things that can happen, you know, you can just
15 be in your home and, you know, cleaning up
16 your bed and there's a big pile of dust, so is
17 that the same as working on the Pile the first
18 day? What difference does it make?

19 Because if you get one little drop of
20 asbestos, then you get that whether you get it
21 on the Pile on the first day or you get it
22 while making your bed, you know, three months
23 later, so it's kind of, I understand from
24 scientifically for us to have all of these
25 categories but working in real-time in what

1 actually happened to people, I think you have
2 to be more open-minded.

3 DR. WARD: And I think we are trying to do
4 that.

5 MS. SIDEL: Oh yeah.

6 MS. FLYNN: I, you know, I have to agree with
7 Steve Cassidy and with Susan Sidel. I mean, a
8 lot of us were involved in the EPA World Trade
9 Center Expert Technical Review Panel where the
10 flaws and inadequacies of all of the
11 government data were, you know, pored over at
12 great length. Unfortunately, the public
13 record of that panel has been removed from the
14 EPA's website and Congressman Nadler's request
15 that it be restored as a resource for this
16 committee and for the public has gone
17 unheeded.

18 But, you know, there have been many, many
19 observations made in that process about the
20 ways in which, for instance, when a monitoring
21 instrument picked up benzene spikes on the
22 Pile, the instrument was shut down and moved
23 to another site.

24 The errors in the, in the asbestos air
25 sampling for lower Manhattan residences that

1 was conducted by ATSDR and the City Health
2 Department were reported by residents who were
3 eyewitnesses to the fact that fans were turned
4 to the wall, that leaf blowers were not turned
5 on. I mean, it almost borders on the level of
6 sampling fraud. So, first of all, they were,
7 you know, we don't have really good sampling
8 data to fully characterize exposures in
9 exposed populations. And second of all --

10 DR. WARD: But didn't I say that? I mean --

11 MS. FLYNN: Yes. No, I just -- I think it
12 really bears reemphasizing and also to -- I
13 know that some people saw this article that I
14 sent in by David Newman, the industrial
15 hygienist with the New York Committee for
16 Occupational Safety and Health, and but I --
17 he said in this article, under the category of
18 exposure assessment:

19 If just one thing is to be learned from the
20 WTC response experience, it should be that an
21 exclusive reliance on environmental sampling
22 data can be misleading and even dangerous.
23 There has been a fundamental disconnect
24 between what the majority of the sampling data
25 would seem to indicate and the breadth of

1 health issues that have arisen. WTC-related
2 illnesses manifested despite reassuring
3 results that came from traditional methods of
4 data collection and assessment. Tens of
5 thousands of WTC responders, area workers, and
6 residents incurred significant and persistent
7 respiratory and other chronic and
8 incapacitating illnesses.

9 And I just want to make one more comment,
10 which is that, you know, not to further
11 complexify (sic) the polling language, but in
12 fact, the Zadroga Act sets a criterion for
13 linkage of illness to World Trade Center
14 substantially likely to have been a
15 significant factor in causing, exacerbating,
16 or contributing to, so is there a way actually
17 to map that language on to the polling
18 language? Because I think we're looking at a
19 real -- I think we're looking at contributing
20 to may get us where many of us feel we need to
21 go much more quickly.

22 DR. WARD: So we can definitely change the
23 language with the poll. I guess I remember at
24 the last meeting there was a little bit of
25 confusion about the criteria for listing

1 something as a World Trade Center-related
2 condition versus the criteria for determining
3 that a particular person's illness was World
4 Trade Center-related. So I don't know if the
5 language that you quoted was -- which one that
6 was. I don't know if it matters, but I think
7 we can certainly change this.

8 I think it really -- what I was -- what we
9 were trying to do is come up with a way to
10 express it where we can understand the
11 diversity of opinions among the group so that
12 we can figure out how we can have a more
13 productive discussion tomorrow. Whether the,
14 you know, if we have general agreement on the
15 overall issue of the potential for
16 carcinogenicity, then we can move on and
17 discuss other things. If not, we need to
18 stick on that point until we understand why
19 different people have different views.

20 DR. HARRISON: Thank you. I wanted to say
21 something else, but I wanted to thank you
22 because I am going to change what I was going
23 to say, I think, because I was not aware that
24 there was the language.

25 And I would ask, maybe, if we could clarify

1 that point because I think, at least in terms
2 of my thinking about whether or how or what we
3 would recommend as a committee, if we need to
4 use certain criteria that is legislatively
5 mandated, I think it's very -- it's
6 significant, pardon the pun.

7 So, if we could just clarify that because
8 there are -- because it actually ties in with
9 the comment that I was going to make. I think
10 there's all sorts of perspectives on how to
11 come to a recommendation in terms of cancer
12 causation.

13 There's the individual patient that some of
14 us, including myself, bring to that
15 perspective when I see an individual in my
16 office with an occupational or environmental
17 cancer, what criteria do I use. There's
18 workers compensation criteria. There are
19 civil litigation criteria. There are cancer
20 presumption law criteria. There are many
21 different frameworks that I personally am
22 familiar with and bring to this discussion.
23 If there are other specific criteria that in
24 the legislation that directs us to consider,
25 then I think we should at least understand

1 what that is and come to whatever straw poll
2 with a reasonably common set of understanding
3 so that -- and this is my comment -- it's sort
4 of agreeing with Steve. It's that if you do a
5 straw poll before we have some common
6 framework may just give us, you know, 15
7 different ideas about what we are voting on
8 but not a common set of criteria to guide our
9 vote.

10 DR. QUINT: Yet another frame is a public
11 health frame and the prevention frame that I
12 come from and also the toxicology frame. I
13 just wanted to tie some of this back to Liz's
14 presentation where she talked about mechanisms
15 because one thing to consider, when she talked
16 about mutations is one -- a lot of these
17 carcinogens are thought to have no threshold,
18 so that when you're talking about amount of
19 the carcinogen or substance that the person
20 was exposed to, it's thought to be linear, so
21 it's going through zero, so any amount could
22 trigger a carcinogenic response.

23 Of course, you know, normally we talk about
24 some risk above background, but to do that,
25 you have to know the potency of the carcinogen

1 plus you actually need to know exposure
2 information and something about the exposure
3 profile: how many days a week, how many
4 years, et cetera, that the person was exposed
5 to it; and we don't have those data.
6 So and the -- there's an article in our file,
7 the folder, Guyton, et al, in Mutation
8 Research which is very compelling because it
9 talks about these carcinogens operating
10 through many modes of action, so it's not just
11 one. It's not just that they cause a
12 mutation. They can act on, you know,
13 promotion and different aspects of the
14 carcinogenic process.
15 So read by my count have 72 carcinogens in the
16 dust, at least the ones that NIOSH listed.
17 Some of these are human. Some of these are
18 animal, so I think, you know, we have to keep
19 all of these things in mind when we talk about
20 biological plausibility.
21 There are a number of in vivo and in vitro
22 articles where people have actually
23 demonstrated with very short exposures, you
24 know, a triggering, mostly the carcinogens
25 that act on an inflammatory process, but, you

1 know, have initiated a process that ends up,
2 you know, that goes through all of the steps
3 and so -- and in very short time periods, some
4 acute and some sub-chronic exposures.

5 Again, they're in mice, and they're in human
6 epithelial cells, but I think all of this
7 enriches our understanding of the mechanisms
8 of carcinogenesis and argues that this is a
9 very complex process when you add, you know,
10 high exposures, very high exposures with a
11 multitude of carcinogens, you add to that
12 complexity.

13 And also ingestion. You can't forget about
14 the fact that some of the exposures probably
15 occurred through ingestion when you have dust
16 on surfaces, especially in offices and homes,
17 you probably have added to that probably also
18 with the firefighters as well, given the
19 amount of contamination on their uniforms. So
20 it's not just the air levels. It's a, you
21 know, very rich mix of information that we
22 have to consider.

23 MS. SIDEL: Just in terms of ingestion, my
24 supply tent was right on the Pile and we were
25 serving coffee and food and all sorts of

1 things, so I'm sure that things were flying in
2 there.

3 DR. WARD: So are you -- oh, Steve.

4 DR. MARKOWITZ: I just want to follow up on
5 what Dr. Quint was saying. So we don't have a
6 lot of experience with people with short
7 exposures and long-term follow-up and cancer
8 in particular, so could you just discuss a
9 little further what experience there is with
10 animals about certain carcinogens with acute
11 or a very short term exposures subsequently
12 relating to cancer?

13 THE COURT REPORTER: Can I say something real
14 quick? If you'll get that microphone real
15 close to your mouth it helps me a lot. I will
16 appreciate it. Thank you.

17 DR. QUINT: I agree with you. Dr. Markowitz
18 said that there isn't a lot of data. I was
19 actually looking for some dose rate data in
20 animals to sort of understand better whether
21 or not we had those models, but there is a
22 paper by Beaver et al that -- let's see, I
23 have it right in front of me here. And
24 actually, she was looking at the exposure to
25 chromium and looking at lung inflammation and

1 injury and then a proliferative -- or from
2 repetitive exposures.

3 And I think in that situation, she was able to
4 expose one kind and then get a response.

5 There's also some information where people are
6 looking now for other than animal models, and
7 so the Hammer Institute had a study where they
8 actually had a training set of carcinogens,

9 NTP, and exposed after 90 days and was able to

10 -- they looked for a marker which was a -- it
11 was a gene expression biomarker, and they were
12 able to see that within 90 days. I think

13 other people have seen it within 24 days, so
14 they're looking at different -- they're not
15 looking at the cancers, but they're looking at
16 markers for carcinogenicity, very specific.

17 There's the other study that I mentioned was
18 the -- a study in human epithelial cells, and
19 I have -- in that study, they were looking, I
20 think, as short a period as 24 hours or maybe
21 shorter than that, and they were looking at --

22 they compared both silica, crystalline silica
23 and amorphous silica and were able to get a
24 difference again in the whole process, you
25 know, leading that was carcinogenic-like

1 process.

2 So, no, animal models, I don't know of any in
3 the regular bioassay models that would mimic -
4 - that we could look at with this.

5 DR. WARD: There's also a lot of data on the
6 cancer patients who were treated with
7 radiation and chemotherapy, and there's very
8 good data on their development of second
9 neoplasms, and in some cases, you will, you
10 know, there's enough data, let's say if
11 someone -- there's a lot of data, for example,
12 on young women treated for Hodgkin lymphoma
13 with high-dose radiation to the chest who
14 subsequently developed breast cancer.
15 So you could look at age and dose if that's --
16 but those are -- those agents are very strong
17 carcinogens, but it is a very rich resource if
18 you're into understanding how relatively
19 short-term high exposures can result in
20 carcinogenic effects, but...

21 Sorry.

22 DR. MIDDENDORF: That's okay.

23 DR. WARD: I keep forgetting about this.

24 DR. TALASKA: There are a number of studies
25 that were done by intratracheal lavage of PAHs

1 that were single-dose were able to bring lung
2 tumors, particularly in strains of mice that
3 were relatively sensitive, so there is --
4 there are data. I can't think of the
5 citations off the top of my head where lung
6 lavage of PAHs, benzo[a]pyrene particularly,
7 has led to a, lung tumors in animals from a
8 single dose, a single heavy administration of
9 a material in liquid -- in corn oil or another
10 vehicle.

11 DR. WARD: Yes, again, I think the other thing
12 to keep in mind is what I mentioned in the
13 presentation that for some of these exposures,
14 they -- if there's a long residence time in
15 the lung and thoracic lymph nodes, a very
16 heavy short-term exposure can result in a
17 long-term dose. And so -- and I think we have
18 some evidence of that in some populations.

19 Okay, so any further discussion before we turn
20 to John Dement's presentation on asbestos?

21 Excuse me? Oh, sorry. Folks on the phone,

22 any further comments before we move into

23 John's presentation? Hearing none, John,

24 would you like to start with your

25 presentation? Well, Paul will queue up your

1 slides and let you know when they're ready.

2 **ASBESTOS AND WTC**

3 DR. DEMENT: Okay, very good. Thank you and
4 my apologies for not being able to be at the
5 meeting today.

6 DR. MIDDENDORF: They're ready any time, John.

7 DR. DEMENT: Okay, just move on to the second
8 slide. I'm going to talk about the dust
9 exposure, so there's clearly the type of dust
10 cloud presented in this photograph is a major
11 high-level exposure to a mixture of things
12 that we have already discussed today.

13 Next slide. There were no measurements done,
14 as we have already discussed, of
15 concentrations in the initial cloud. I think
16 Paul Liroy and some others have estimated that
17 the concentrations were likely in excess of
18 100,000 micrograms per cubic meter, 100
19 milligrams per cubic meter.

20 And I've sampled some industrial operation as
21 a hygienist where dust levels were
22 consistently in the neighborhood of 20 to 30
23 milligrams per cubic meter, not as high as
24 this. So I think this estimation is probably
25 a reasonable estimation, maybe on the low side

1 for at least the initial dust cloud.

2 Lioy described what he considered, and I think
3 is a reasonable consideration, five specific
4 post studies on 911 exposure categories.

5 Go to the next slide. And clearly the highest
6 exposed were those there during the initial
7 collapse and the days that occurred

8 afterwards. I understand there was a rain
9 event like around the third day, which helped
10 to dampen at least some of the dust exposures,

11 but I think the scenario is something like
12 this: We have high-level exposures initially,

13 and then we have continued exposures to the
14 individuals who were doing the recovery and
15 cleanup longer term, and also exposures to a
16 much more mixed of (indiscernible) and fires
17 and materials in that.

18 Let's go to the relative -- next slide,

19 please. One of the relatives to dust exposure
20 is (indiscernible) based on the plume depicted
21 in this slide. I think clearly the first day,

22 extremely high exposure, followed by lower-

23 level exposures during some of the recovery

24 operations; however, if I could point out

25 here, there were no dust measurements actually

1 made on this first day. So these are
2 reasonably speculative.

3 I am going to talk about asbestos, and go to
4 the next slide please. And I am going to talk
5 about some of the measurements that were made.
6 First, I wanted to talk about the methods that
7 have been used for measuring asbestos
8 exposures, both historically and currently.

9 On the list on here is an old midget impinger
10 method developed by the U.S. Public Health
11 Service in the 1920s. It's been used, really,
12 for exposure measurement in occupational
13 settings for dust exposures up until about the
14 mid-1960s. I mentioned that largely because
15 the old dust measurements and the basis for a
16 lot of the risk assessment for asbestos are
17 based on the old impinger method.

18 First of all, it was a method that didn't
19 collect fibers very efficiently. Secondly,
20 the exposure method actually counted all
21 particles, not just fibers in the dust and it
22 did it at a low power using low power optical
23 microscopes.

24 So there's some -- excuse me -- some severe
25 limitations with regard to retrospective

1 exposure assessments even in the occupational
2 environment. The current method used has been
3 used since about the 1960s. It's called phase
4 contrast microscopy. Basically the samples
5 are collected on a filter, membrane filter,
6 and the particles counted by an optical
7 microscope that has a special feature which
8 enhances contrast called a phase enhancer.
9 But still, it's relatively low magnification,
10 400 times.

11 There are certain limitations to this method.
12 First of all, the cause of limitation with
13 regard to being able to count short fibers.
14 Only fibers longer than five micrometers are
15 counted. Secondly, even if a fiber were
16 longer than five micrometers, this counting
17 system -- the microscope has no resolution or
18 ability to actually see small diameter fibers.
19 So you could have very long fibers that were
20 small in diameter and not be detected.

21 Nonetheless, it's used as part of the OSHA,
22 current OSHA standard, and it's the basis of a
23 lot of the risk assessments. And I think it's
24 -- the use of the phase contrast microscope
25 has actually enhanced some misconceptions

1 about the nature of exposures and what's
2 important. That is, only long fibers or
3 fibers longer than five micrometers -- I'm
4 going to have more to say about this later.

5 Moving on to scanning and transmission
6 electron microscopy. Scanning microscopy is
7 better than phase contrast, but still not
8 capable of seeing the very small diameter
9 fibers in an asbestos dust cloud.

10 The most useful method is transmission
11 electron microscopy, and some of the measures
12 of the World Trade Center exposures were done
13 by TEM. There are different techniques that
14 are used for expressing the concentrations.
15 Some express structures per centimeter of
16 surface. Some were expressed as structures
17 for -- as a dust concentration measurement per
18 cubic centimeter of air samples.

19 The limitation here is the fact that when you
20 look at samples by transmission electron
21 microscopy, you look at a very small portion
22 of the dust cloud, and it's very expensive.

23 A little bit about the measurements that were
24 done. The range of asbestos, primarily
25 chrysotile, looks like from a less than one

1 percent up to about three percent of the mass.
2 And with most fibers being less than five
3 micrometers in length, which you would expect
4 given the length -- given the nature of the
5 collapse, the pulverizing of material.

6 There's more to say about the less than five
7 micrometer criteria as well because even in
8 asbestos-exposed occupational cohorts, the
9 majority of exposure is to fibers that are
10 less than five micrometers in length,
11 typically 90 percent of actual.

12 Again, no measurements were made of chrysotile
13 during the extraordinary high dust cloud
14 exposure. There was a range of exposure
15 measurements done later and reported in the
16 literature, some in peer reviewed
17 publications, some in -- just in reports.
18 Most of these seem to show short-term
19 exposures of not in excess of established
20 criteria; however, there are lots of
21 limitations of these as we've discussed
22 already. One is reading the samples would be
23 the preferable method for looking at exposures
24 to individuals on the Pile.

25 NIOSH did some sampling on these, used PCM and

1 looked at some of the samples by transmission
2 electron microscopy, and in general, when you
3 look at the samples by TEM, the concentrations
4 didn't exceed the OSHA PEL of 0.1 fibers per
5 cubic centimeter of air. Again, that's fibers
6 longer than five micrometers.

7 Realizing of course that the majority of
8 fibers in the study are less than five
9 micrometers in length. I think there is a
10 disjoint, and I think Liz pointed that out.
11 This dust cloud was extremely high in dust
12 levels, certainly initially. No measurements,
13 again, but we would expect that in that dust
14 cloud, given a concentration of one percent or
15 even much, much less, that the asbestos
16 exposures to total fiber concentration would
17 be very high.

18 I'm going to talk little bit about the types
19 of regulated asbestos because many of the risk
20 assessments have just considered asbestos as
21 one group of materials; that's a list of them.
22 We're dealing largely with chrysotile here
23 which was in the towers.

24 I am going to say there may not be amphiboles
25 in there. I had the opportunity of being in

1 the World Trade Center a number of years
2 before 9/11, and I think there might have been
3 at least some amphiboles in the building as
4 well at some point in time.

5 Liz has already pointed out, I think, that
6 asbestos is considered a carcinogen by both
7 IARC and the National Toxicology Program.
8 That includes chrysotile, certainly with
9 regard to lung cancer mesothelioma. There's
10 no question with regard to the
11 carcinogenicity.

12 IARC also determined that there was sufficient
13 evidence in human studies for cancers of the
14 larynx and ovaries and limited evidence for
15 colorectal and in the pharynx and stomach.
16 And there have been a number of reviews of
17 cancers at sites other than the lungs for
18 asbestos.

19 I think this determination by IARC is
20 reasonably consistent with the data that
21 exists, largely with regard to cancers of the
22 GI system. Studies that show an excess risk
23 of about two for lung cancer tend to show an
24 increase, not a two, but an increased risk for
25 GI cancer.

1 I'm going to talk a bit about the risk
2 assessments that we have for asbestos. Nearly
3 all of the risk assessments are based on
4 populations occupationally exposed. Again, as
5 discussed before, the measurement method is
6 this phase contrast microscopy where the
7 fibers longer than five micrometers in length
8 are counted.

9 The typical metric is cumulative exposure
10 expressed as the product of duration and
11 concentration measured in fiber-years. I want
12 to point this out because a lot of the data
13 upon which risk assessments are made is really
14 occupational groups with short exposures which
15 are relative to high concentration, including
16 the studies that our group has done of
17 chrysotile-exposed textile workers.

18 Many of these workers had exposures of just a
19 few months and nonetheless showed increased
20 risk. Most of the models, including our own,
21 were no-threshold models; that has been
22 discussed already today.

23 They seem to fit best to the actual data. And
24 lastly, a point that needs to be emphasized is
25 that there's no scientifically justified

1 threshold for asbestos-related cancers, none
2 that's been established in the literature by
3 recent studies.

4 Here are the limitations of the risk
5 assessment, moving to the next slide.

6 Historical measurements, as I said before, a
7 lot of them were based on the old impinger
8 method and unless you had some data to make a
9 statistical conversion between the old method
10 and new method, there's lots of
11 misclassification in the data. And in most
12 cases, in these types of studies, that tends
13 to actually dampen the exposure-response
14 relationship. So your effect is likely
15 greater than you are actually showing in your
16 data.

17 Again, the risk assessments were based on the
18 phase contrast method wherein only a fraction,
19 and typically less than ten percent of the
20 actual airborne aerosol was actually measured.
21 And as I said before, that's because of the
22 diameter limitation of the PCM method and
23 because of the decision to count only fibers
24 longer than five micrometers. That decision
25 is really not based on the decision that short

1 fibers are without risk.

2 It's based on the fact that a practical method
3 hasn't been developed for measuring exposures
4 and enforcing standards. And NIOSH, in its
5 1972 criteria document for asbestos pointed
6 out that the reason for the five micrometer
7 cut was for reproducibility of the PCM count.
8 Lastly, mesotheliomas are not well captured in
9 a lot of the mortality data that's been
10 published at least through 1999. There was no
11 code for mesothelioma specifically. Only in
12 ICD-10 do we have a specific code for
13 mesothelioma, so a lot of the mortality
14 studies, including our own, looks at things
15 like cancers of the pleura and assumes that
16 those are mesotheliomas. And that's a
17 reasonable assumption in most cases but likely
18 does not capture well in other cases.

19 Next slide. I wanted to drive home the notion
20 about what portion of fibers are actually
21 counted by phase contrast microscopy. This is
22 actually a slide from some of our data from a
23 textile operation where they're using very
24 long fibers, the best grade chrysotile. And
25 even in textiles, if you look at this

1 distribution of diameter to length, you see
2 that the vast majority of the fibers are short
3 and thin. So that's the nature of exposures,
4 even occupational.

5 Next slide. I wanted, last, to point out two
6 studies that have been published subsequent to
7 the current risk assessments used for the OSHA
8 standard. The two case-controlled studies,
9 and these were for the mesothelioma, one in
10 France and one in Germany, and they are of
11 reasonable size, particularly the France
12 study. And what these studies are showing is
13 that we now have measured excess risk of
14 cumulative exposures that is fiber-years. In
15 the France, study in France, less than one
16 fiber-year.

17 Likewise, in the study in Germany we have an -
18 - about an eight-fold risk for fiber exposures
19 that are less than 0.2 fiber-years. There is
20 a, I think, a legitimate discussion in the
21 literature about the relative ability of
22 chrysotile versus the amphiboles to produce
23 mesothelioma.

24 I think, first of all, there's no question if
25 chrysotile does produce mesothelioma. Whether

1 or not it's less potent than amphiboles is a,
2 I think, a subject for considerable debate.
3 Next slide. Lastly, I want to point out that
4 the OSHA PEL, which is being used as a
5 criterion in some of the assessments of the
6 air samples from the World Trade Center on 0.1
7 fibers per cc as an eight-hour time-weighted
8 average is not without risk. OSHA's risk
9 assessment indicates that at .1 fibers per cc
10 over a working lifetime, there's an excess
11 risk of 3.4 cancers per 1000 workers, and of
12 those 3.4 cancers, about two-thirds of them
13 are lung cancers. The other third are
14 mesothelioma.

15 So, the point is that we don't have a
16 threshold for the cancer-producing effects of
17 asbestos, including chrysotile. It's open for
18 discussion.

19 DR. TALASKA: John, Glenn Talaska. Thank you
20 very much. I've got a couple of questions for
21 you on -- you cleared one up right at the --
22 in your last slide. I wanted to know the
23 relationship between the numbers of lung
24 cancers seen with asbestos exposure documented
25 versus the number of mesotheliomas, and you

1 said the ratio is about two-thirds to one-
2 third.

3 But I also wondered what it was in terms -- if
4 there were any data in terms of latency time
5 relative to those two diseases.

6 DR. DEMENT: Well, I think the latency times
7 are as Glenn just pointed out. Early in the
8 lung cancer, in our own studies, we started to
9 see a pickup in the relative risk, between 10
10 and 15 years and it really starts to escalate
11 after about 20 years.

12 Mesothelioma has what appears to be a longer
13 latency in many cases. The peak of that
14 probably, in most states, hasn't occurred
15 until 30-plus years after a person is exposed.

16 DR. TALASKA: Thank you, and I have one
17 further question. You didn't talk about it.
18 I am only going to mention it briefly in the
19 next presentation, and I hope you will join me
20 in the discussion then of the interaction
21 between things like PAHs and asbestos. Do you
22 want to give a little -- if you had some
23 information you could provide us right away or
24 would you -- we could wait until after my
25 talk, because I am going to just mention it

1 briefly.

2 DR. DEMENT: I'll mention it briefly as well.
3 I think in lung cancer, there's clearly an
4 interaction with PAHs and particularly
5 smoking. The question is whether or not
6 that's a multiplicative additive or less a
7 multiplicative fact, and I think most
8 individuals, it may not be multiplicative but
9 it's more than additive, so there is an
10 interaction there. I guess we can discuss it
11 later.

12 DR. WARD: Other questions or comments for
13 John? John, I -- one question I had was if in
14 the two case-controlled studies with
15 mesothelioma, it was hard for me to
16 conceptualize, you know, how small those units
17 were. Can you help, I mean, can you compare
18 it to like what a typical occupational
19 exposure would be?

20 DR. DEMENT: Well, these levels are, if you
21 look at the fiber-years, most occupational
22 risk assessments are based on a 40 or 45 year
23 lifetime risk, working lifetime risk. So if
24 you take the current OSHA standard of .1
25 fibers per cc over a 45 year working lifetime,

1 that's 4.5 fiber-years.

2 These data, these case-controlled data, are
3 clearly demonstrating excess risk at exposures
4 that, cumulative exposures that are much less
5 than that, which just really adds to the
6 conclusions of the OSHA risk assessment. That
7 is, these are not zero risk standards.

8 The OSHA standard includes lots of work
9 practices in an effort to try to get exposures
10 as far below this .1 fibers per cc as
11 possible. The other thing I like to point out
12 is the occupational cohorts. There are
13 cohorts, including ours as I mentioned before,
14 that do demonstrate excess risk with short-
15 term workers at relatively high levels of
16 exposure, of course.

17 The one that was done in Paterson, New Jersey
18 by Dr. Seidman in Mt. Sinai many years ago
19 demonstrated that individuals who worked down
20 in that plant with one month of exposure
21 producing asbestos, they had a significant
22 excess risk of cancers, including lung cancers
23 and mesothelioma.

24 DR. WARD: John, can you comment on half-life?
25 I mean are the -- I mean, I know that

1 different types or lengths of asbestos would
2 have different residence in the lung, but is
3 there -- I mean, there probably have been
4 studies looking at pathologic specimens of
5 workers exposed to asbestos. I mean, does it
6 tend to stay in the lung for a long time?

7 DR. DEMENT: What it does -- there is some
8 discussion, certainly in the literature with
9 regard to the clearance rates of amphiboles
10 versus chrysotile, and in general I think the
11 amphiboles cleared less quickly than
12 chrysotile.

13 There was a study done at Mount Sinai by Dr.
14 Suzuki, who suggests that the clearing of
15 chrysotile from the lung actually ends up
16 concentrating in the pleura where we actually
17 see mesothelioma in the study.

18 I think the studies that have looked at lung
19 burden are sometimes problematic with regard
20 to chrysotile because of its (indiscernible),
21 and I think some erroneous conclusions have
22 actually been drawn based on lung burden
23 studies when you didn't actually have the
24 estimates of the actual exposures to the
25 individuals.

1 DR. HARRISON: This is Bob Harrison. Steve
2 Cassidy, earlier this morning, earlier this
3 afternoon, sorry -- I'm on West Coast time --
4 suggested that the samples may not have been
5 representative of the type of exposures or
6 type of activities that people had. I wonder,
7 John, if you could comment on that.

8 You said that samples weren't taken, I guess,
9 in the first three days. And then there were
10 lots and lots of samples taken subsequently,
11 but I don't have a clear picture of what
12 people were doing, where those samples were
13 taken, and whether there were other activities
14 where we think exposures were probably higher
15 that were not captured.

16 DR. DEMENT: Well, I don't have a good sense
17 of that either. My sense of the data itself
18 is that most of the personal air sampling that
19 was done was either done by NIOSH or NIOSH
20 contractors through NIOSH. Those were
21 represented in the publication, I think, by
22 (indiscernible) through NIOSH, and in the
23 slide, where we showed (indiscernible)
24 samples.

25 A lot of these were actually taken during the

1 post-cleanup operation, but the extent to
2 which they represent exposures of that group
3 is really not known. I mean, an effort was
4 made to do that, but, you know, I can't, you
5 know, I don't know all of the cache that were
6 not sampled.

7 DR. WARD: Any other questions or comments?
8 Susan.

9 MS. SIDEL: Hi, John. Susan Sidel. Could you
10 just explain again the different measurements
11 that you used that -- you were saying a TEM is
12 the -- is like the finest but it's also really
13 expensive and it's not OSHA standard. So the
14 OSHA standard doesn't pick up the tiniest
15 particles, and what was used at the World
16 Trade Center?

17 DR. DEMENT: The OSHA standard is based on the
18 space contrast method.

19 MS. SIDEL: Right.

20 DR. DEMENT: So it's an optical microscope
21 with a phase -- a phase illuminator or phase
22 shift illuminator, and the problem -- just go
23 back and place yourself in the 1960s. All of
24 the old samples were collected by methods
25 including (inaudible) with a routine sampling

1 method that would first of all actually
2 measure fibers, if not all particles, and
3 measured a reasonable portion of the air
4 samples.

5 So this method was the default method, and it
6 measures, even in the asbestos industry,
7 occupationally, it is really just an index of
8 exposure. It's measuring a small fraction of
9 the air blowing aerosol. Because of the
10 limitations of the counting with regard to
11 length and the resolution with regard to
12 diameter.

13 So, typically, in an occupational setting with
14 chrysotile in particular, because it tends to
15 be more fine, you'd be lucky if you're
16 counting 10 percent. In most cases, you're
17 counting about five percent of the total
18 number of asbestos fibers that are airborne
19 that the workers are actually breathing.

20 If you move on to electron microscopy, it has
21 the ability to look at these particles, but
22 because of the high magnification, you're
23 actually looking at a very small area of the
24 filter, so you have a lot of statistical
25 variability with regard to the count. It was

1 not chosen as the method for routine
2 occupational exposure assessment.

3 MS. SIDEL: So the method that was used in the
4 World Trade Center is the method from the
5 1960s?

6 DR. DEMENT: Sorry, could you repeat?

7 MS. SIDEL: So the method they were using at
8 the World Trade Center was the OSHA standard
9 method that you talked about from the 1960s?

10 DR. DEMENT: No. Yes, most of the samples
11 that were workplace samples. For example, if
12 you look at the slide, 19,000 air samples --

13 MS. SIDEL: Uh-huh.

14 DR. DEMENT: Almost all of those were PCM, so
15 they did not use transmission electron
16 microscopy. So it's trying to measure these
17 exposures against an OSHA standard. The NIOSH
18 sampling used PCM, but they did -- didn't look
19 at the ones that were in excess of the .1
20 fibers per cubic centimeter and looked at
21 those by TEM. Samples which were mostly
22 structures per millimeters squared filter area
23 were TEM.

24 MS. SIDEL: Thank you.

25 MS. HUGHES: Hi. I just want to remind

1 people, as a resident that lived one block
2 away, the chaos that was there for a very long
3 period of time, there was no electricity. So
4 if you're going to do sampling or testing and
5 there's no electricity, one of the concerns
6 that some of the testers had was it could be
7 done on a generator, and then you had to
8 determine what kind of generator.

9 Would you be using diesel fuel, or would you
10 be using a battery, and then where you would
11 get that. So there was electricity on the
12 east side of Broadway but not the west side of
13 Broadway, and so when people are talking about
14 the proximity of the testing, it took some
15 time to actually get the machinery into place
16 to actually do the testing.

17 And then one of the issues that has been
18 argued about over the years was clogged
19 samples, so the filters were clogged if there
20 was a lot of material that was actually picked
21 up. So I just wanted to remind people what it
22 was like early on. Thanks.

23 DR. DEMENT: Those are good points to make. I
24 think given a relatively low percentage-wise
25 of asbestos in this material and the high

1 concentrations of dust, one of the issues with
2 regard to asbestos sampling is trying to
3 optimize the ability to count it, and when you
4 run a filter for a period of time,
5 accumulation of dust on the filter can
6 actually obscure the PCM count.

7 DR. HARRISON: This is Bob Harrison. I just
8 wanted to make two points. I think both of
9 them are probably obvious, but I think for the
10 record, it's worth stating. One is that I
11 think there's evidence that respiratory
12 protection was not available, consistently
13 used, and would not have afforded, in any
14 event, protection against inhalation of
15 potentially carcinogenic asbestos fibers.

16 I don't -- I'm not sure that there would be
17 any disagreement about that point, but I think
18 it's worth noting and if there's any, you
19 know, any additional comment, we need to make
20 that.

21 The second is that based on the lung disease
22 that we've seen from other lines of evidence,
23 (indiscernible) airways tends to show
24 (indiscernible) lung diseases. I think we can
25 use that as qualitative evidence that indeed

1 inhalation of particles and fibers and smoke,
2 et cetera, did occur.

3 I don't think we can make any correlation
4 between those clinical effects and the dose of
5 asbestos, but I think just qualitatively, we
6 know that this population had inhalation
7 exposure, and I just think it's important to
8 point that out as well.

9 DR. MIDDENDORF: John, this is Paul. I just
10 want to ask if you would take a minute or so
11 and address the issue of potency related to
12 length of asbestos fibers.

13 DR. DEMENT: Well, I think, Paul, the issue of
14 potency with regard to length, it really comes
15 from some animal data. Now if humans are
16 exposed to the whole spectrum of fibers, and
17 so when I studied my textile workers, they're
18 exposed to the whole dust cloud irrespective
19 of how I choose to measure it.

20 Some of the animal studies suggest longer
21 fibers are more carcinogenic, and those
22 studies come from some inhalation, but mostly
23 studies that are implantation are injection
24 studies, some of the early studies from Merle
25 Stanton at the National Cancer Institute, for

1 example, and Dr. Hoch (ph) in Germany.

2 So with regard to cancer, I think
3 longer/thinner may be more carcinogenic, but
4 in the exposed aerosol, even if you consider
5 an asbestos textile, the longer/thinner
6 comprise a very small portion of the airborne
7 exposure.

8 So I think the -- in terms of the actual
9 effect of short fibers in that they greatly
10 outnumber the long thin ones, even if fiber
11 for fiber, they were a fraction -- had a
12 fraction of the carcinogenic potential, I
13 think the data doesn't support leaving those
14 out with regard to risk assessment. We just
15 completed a series of studies in the plants
16 that we've looked at for many years in South
17 Carolina and in North Carolina, and we did
18 these in collaboration with NIOSH where we had
19 the ability to go back and look at some of
20 those old filters in the 1960s and to try to
21 estimate a sort of size specific exposure
22 measurement for these workers in these two
23 cohorts and try to relate that to risk.
24 And when we did that, we found that all of the
25 size categories by length and diameter

1 correlated and predicted lung cancer risk.

2 It's -- the longer, thinner fibers, when you
3 look at them had a slightly greater impact;
4 but nonetheless, all sizes that we were able
5 to measure, including the short thin ones,
6 impacted lung cancer.

7 DR. WARD: Any other questions or comments on
8 this presentation? Thank you very much, John.
9 We hope you can stay on for some more of the
10 discussion. We appreciate you coming.

11 DR. DEMENT: I'll plan on staying on. Thank
12 you.

13 **PAHs AND WTC**

14 DR. TALASKA: Okay, are we ready? How does
15 that sound? Good? Everybody okay? Okay.
16 Well, I wanted to begin by making a statement
17 about how being able to look at these data in
18 detail, really it changed my mind about
19 something about the exposure with the -- of
20 the first responders at the Ground Zero site.
21 When I, as a scientist, and as a regular
22 scientist with an interest in the area, but
23 not an acute one, I looked at the abstracts.
24 I looked at some of the tables, certainly of
25 the ones with biological monitoring because

1 that's my field.

2 And -- but I didn't look at the papers really
3 hard, and the opportunity that I got today to
4 look at them -- today -- in the past two
5 weeks, at least, and certainly since being on
6 the committee has given me a somewhat
7 different -- considerably different
8 perspective than I've had to begin with, and I
9 will begin with this.

10 What I'm going to talk about today are the
11 polycyclic aromatic compounds. These are the
12 materials that are formed by the burning of
13 any material as a fraction of the total mass
14 of the stuff that's burned. Most of the stuff
15 goes to carbon dioxide, but if there's not
16 sufficient oxygen to go to complete oxidation
17 of it, then these benzene rings fuse and form
18 large plate-like structures that I give you
19 three examples here.

20 These are materials that -- from any kind of
21 burning. I'll show you some pictures. PAHs
22 are very lipophilic materials. They're well
23 absorbed from both the lungs and the skin when
24 they're contacted and from the GI tract,
25 although there is a difference with the GI

1 tract relative to these compounds that I'll
2 get at later.

3 Just some examples from the occupational world
4 first. You can see from here -- there it is -
5 - that the upper left panel shows a coke oven.
6 This -- the worker here is a topside coke oven
7 worker -- these two workers. One of them is
8 more obscured by the smoke than the others.
9 These are occupational exposures where we have
10 both the knowledge of what the internal dose
11 was for these individuals and the lung cancer
12 risk, which is at excess. These people are in
13 the worst possible situation because you're
14 trying to make coke, not Pepsi-related coke,
15 but coke which is used in steelmaking out of
16 coal.

17 So it's burned in the absence of oxygen or
18 almost the absence of oxygen and forms a dense
19 smoke which escapes from the machine. It's a
20 very large structure. The right-hand panel is
21 a foundry. And you can see, again, the hot
22 metals are producing smokes which can be seen.
23 The lower right-hand panel is an aluminum
24 manufacturing site. At this slate, they're
25 pouring.

1 The left one is extremely interesting from
2 several points of view. One is it's a food
3 product. Our PAHs are in many of the foods
4 that we like. Barbecue, smoked foods contain
5 PAHs from the prioritization of the materials,
6 and we eat them.

7 But also look at this here. As you can see
8 from closer examination of the walls of the
9 smokehouse that this guy is in smoking fish,
10 that the whole structure is coated with a tar-
11 like substance. And those are -- that is
12 often high in the -- very high in the PAHs.
13 Other examples are shown here. This slide
14 shows an asphalt operation that we've all
15 smelled. The materials that are coming off
16 the gassing of the asphalt as we, you know,
17 our body -- I think everyone uses orange
18 barrels. And so the workers are exposed
19 there.

20 One of the real advantages of the studies that
21 have been done very much by NIOSH but with
22 other players as well is that often times they
23 will take area samples of areas near or around
24 a -- some of these operations and then conduct
25 personal samples at the same time. And that

1 becomes important to us.

2 In the right-hand panel is the classic PAH
3 exposure that causes lung cancer in cigarette
4 smokers. Seven to ten-fold excess risk,
5 depending on how many packs are smoked. It
6 goes up with a various dose response that most
7 of the toxicology is envious of, but it's from
8 a very sad point of view that this is the
9 major carcinogenic material in the United
10 States and the world for causing lung cancer.
11 PAHs are also formed with the burning of any
12 material, so the nasty smell that you get when
13 the smoke comes your way at the campfire
14 contains some of those materials and that's
15 the stuff that stays on your clothes the next
16 day when you realize that, you know, those
17 were in a bar or where there was smoke.
18 The lower right-hand panel, of course, shows a
19 more recent disaster caused by -- during the
20 blowout last year of the oil rig in the Gulf,
21 the Deep Water Horizon. And you can see --
22 and this is important from -- for our
23 discussion because you can see two things.
24 One is that here is where the closest you can
25 get to this thing to do any sampling at all is

1 the distance, several boat lengths between the
2 fire and the -- and the source of the burning
3 itself.

4 And then you can also see the huge difference,
5 if you collected a sample here, what would be
6 the exposure level relative to what it would
7 be if someone was at or near the plume? I'm
8 not making direct comparisons, but keep this
9 model in your mind is what I'm saying there.

10 And now we have the World Trade Center and
11 slides that I have -- a couple of slides just
12 to illustrate things about the smoke. Here we
13 have a burning smoke which is -- probably has
14 PAHs in it, almost certainly, and then the
15 more general smoke that occurred, I believe,
16 right after the collapse where the -- probably
17 a multitude of materials in this one.

18 Also important here is that at this point you
19 can see there are civilians inside of this
20 where they -- where the work is actually being
21 done. Now, I'm not sure, and I have to tell
22 you I don't know as well where the monitors
23 were put at Ground Zero relative to the work
24 zone.

25 And -- but that's extremely important. Even

1 at this point, you can see your, you know, the
2 smoke is going up. Oh, that was the other
3 thing with this one. I'll go back a minute.
4 The smoke is rising here very rapidly.

5 Persons that are in the plume are being
6 heavily exposed, but persons very, just to the
7 outside of it, outside of the convection
8 currents that are occurring, are not being
9 exposed to the same levels. Nor would any
10 monitors that are placed in that area be
11 exposed to the same level.

12 Okay. PAH exposures are associated with lung
13 cancer in tobacco smokers. It's thought that
14 70 percent of the lung cancer in the United
15 States and the world is due to tobacco
16 smoking. Coke oven workers are also at
17 increased risk. Aluminum smelter workers are.
18 And the classical exposure to -- of soots,
19 dermal exposure on the scrotum in chimney
20 sweeps was investigated by Percivall Pott in
21 1776 and associated with the soots that were -
22 - people, kids mostly, who were exposed to
23 that by actually being run through the
24 chimneys at the time.

25 The PAHs are absorbed by the body and they are

1 metabolized to compounds by the body that
2 combine to DNA. So PAHs themselves are not
3 carcinogenic. It's the PAH metabolites that
4 are carcinogenic, bind to DNA, and cause
5 mutations that initiate the carcinogenic
6 process. So it is biologically plausible that
7 PAH can cause cancer if there is sufficient
8 exposure.

9 What are the sources of combustion materials
10 at the World Trade Center? This has been
11 reviewed in a NIOSH document, and I'm just
12 showing it for you.

13 There was approximately 90,000 liters of jet
14 fuel, 500,000 liters of transformer oil,
15 380,000 liters of diesel and heating oil, and
16 approximately, although no one knows for sure,
17 the same amount of gasoline which was burned
18 in the parking structures when the towers
19 collapsed and over the next several days as
20 those cars heated up and exploded or were
21 demolished and then the gasoline leaked all
22 over the place and then burned.

23 Area samples were collected and for PAHs
24 specifically, not for dust in particular, but
25 for PAHs in particular, were collected at the

1 fence line beginning on 9/16 through 9/23/01.
2 There were no personal samples taken at this
3 time by these investigators. So the first
4 samples seem to be five days after the
5 exposure. There were biomarker samples
6 collected once on October 1st, approximately,
7 in a study that was reported by Edelman et al
8 in 2003.

9 But I think it's also interesting, and I'm
10 going to bring up the set of studies that I
11 found in the Butt et al 2004, a Canadian group
12 who looked at the window films and extracted
13 the materials from the films of windows at
14 various places in New York City and found
15 considerably different levels of PAHs on them
16 than were collected in the air samples.

17 So these are the data of Pleil et al at the
18 fence line, and again, area samples. You can
19 see many samples were collected throughout.
20 Samples were collected at the perimeter of
21 Ground Zero, not in the work area, but at the
22 perimeter and again, no samples for the first
23 five days.

24 They were also collected distally at Broadway,
25 so away from the site. And one of the things

1 that you can see clearly is that these two
2 exposures have parallel curves. They run
3 together down here, but they're parallel
4 pretty much out here. So we have a difference
5 between the two of them by at least a factor
6 of two because based upon the distance.
7 So -- but again, they were area samples,
8 stationary samples collected not following any
9 particular worker, not following any
10 particular activity at all, but sitting at the
11 fence line, some distance from where the
12 activities were being taken -- taking place.
13 So all of these samples are -- were air
14 measurements and estimates based on area
15 samples collected at the fence line, and these
16 types of samples typically underestimate
17 worker exposure and the differences can be
18 anywhere from three- to 40-fold, that if you
19 take an area sample at a periphery, depending
20 on how far away it is from the active sites of
21 the workers, it generally is known to
22 underestimate the exposure.
23 Now, that difference can be even greater than
24 40-fold, but it can be less than 40-fold as
25 well, and the way that it can be less than 40-

1 fold is if the study design uses an area
2 sample to capture the worst case. So many
3 times in my career, I've stationed an area
4 sample in the worst possible exposure place
5 where there are no workers, but to capture the
6 worst-case scenario to see -- and the idea
7 being if there's no problem at the absolute
8 worst designed place, then there might not be
9 a problem where the workers are.

10 But one has to consciously design their study
11 to do that to be able to catch a worst-case
12 scenario, and I don't believe that was done in
13 the studies that were collected. Secondly
14 -- so we have a difference here that could be
15 fairly large. Secondly, only the PAHs that
16 were in the particulate phase were counted
17 because they captured the 2.5 micron samples,
18 extracted those samples.

19 There's also PAHs in the vapor phase. PAHs,
20 if they're heated, turn into a vapor, like
21 steam, and then that steam rises into the air.
22 And that is -- sometimes it binds to particles
23 and it does bind to particles, but some of it
24 stays in the vapor phase as well.

25 And depending on the type of study -- in

1 Burstyn et al there was -- they found 10 times
2 more PAHs found in the vapor phase than
3 asphalt workers, but other workers have seen
4 things much lower.

5 So they have seen 10 times more in this one
6 study, but Quinlan et al, for example, in coal
7 liquefaction workers saw that the amount that
8 was in the particulate, bound particulate, was
9 about equal to what was found in the vapor
10 phase. And there are estimates all over the
11 place between those extremes.

12 Okay. So what effects weren't measured?

13 Well, the first question is what is the impact
14 of being in a plume and how much more would
15 that be, and how much greater, and again, I
16 refer you back to the picture for the Deep
17 Water Horizon.

18 If you're working right above the smoke as
19 opposed to being away from it at the
20 periphery, then the -- what would be the
21 impact? And I have -- unfortunately, I wasn't
22 there, and I can't tell you.

23 What is the effect of exercise and exertion,
24 and I'll show you a slide about how important
25 that can be. But if somebody is working hard,

1 they are breathing hard and they are breathing
2 several times more than what the, on average,
3 if I am working really hard riding a bicycle
4 or jogging, you know, the worst place to jog
5 is along city streets.

6 Fortunately, the lead's out of gasoline but,
7 you know, the worst place to jog is around
8 there because you are breathing several times
9 more and that means you are breathing more of
10 this material into your lungs where they can
11 be collected.

12 So that's an impact that one might want to
13 consider, especially if different groups of
14 people were working harder. From what I can
15 gather, and I think in the paper, in the Pleil
16 et al paper, they estimate that -- the purpose
17 of their sampling was to look at some general
18 environmental effects. They weren't looking
19 for what was happening to the workers at
20 Ground Zero, okay, so -- and they made no
21 attempt to capture the peaks or assess exposed
22 worker exposure, and they stated specifically
23 that exposure to the workers at the site could
24 be quote, much higher, end quote.

25 So there is a big weakness with the best PAH

1 studies that were done at the site, and now --
2 oh, yeah, but here is something that I believe
3 is illuminating as I was going through the
4 voluminous literature that was provided us.
5 Butt et al did a series of studies where they
6 washed windows with solvents, and they washed
7 the windows to be able to extract the PAHs and
8 other materials. They were looking for PAHs
9 on them, okay? And what they saw was that
10 there were different zones and -- as you might
11 expect.

12 So within one kilometer -- they are Canadian
13 after all -- which is 6/10 of a mile, the
14 average was 77,100 nanograms per square meter.
15 We were seeing in the other study, in the
16 Pleil et al study that they were talking about
17 35 nanograms per cubic meter, so a meter is
18 three feet approximately by three feet by -- a
19 cubic meter is three feet by three feet by
20 three feet. A square meter is three feet by
21 three feet, but on average, Butt et al were
22 seeing on these window films which admittedly
23 collected samples for several days, they -- I
24 forget the day that they collected them on --
25 they were considerably higher, thousands of

1 times higher.

2 In fact, downwind sites within one kilometer
3 averaged 130,000 nanograms per square meter.
4 Upwind sites were much lower, averaged 18,500,
5 still within a kilometer. Upwind sites that
6 were greater than two kilometers away averaged
7 6000, and this might be considered the
8 background for New York City windows, okay?
9 More than two kilometers away, and upwind, so
10 the wind from the site probably wasn't blowing
11 very often on these windows.

12 So you can see the types, now, you know, you
13 can't use this for exposure estimates,
14 obviously, but these are windows that may or
15 may not have been in the major plumes at all.
16 By luck, they sampled these, and I don't
17 believe they had any selection other than they
18 had access to the buildings. So I thought
19 this, this was illuminating to me.

20 Here's some of the data about work rate. So,
21 if you are working, light work is what we
22 consider for most of our standards where the
23 work load in watts is about 50 watts that the
24 alveolar vent -- so, at rest, the people that
25 are in this room are breathing in about five

1 liters of air per minute, but someone who is
2 working very hard can breathe seven times
3 that. So they bring in seven times the amount
4 of air. They pump the blood around much more
5 efficiently. And so you can see the exposure
6 metrics can give you another twofold over that
7 if you're worried about heavy work as opposed
8 to light work in terms of the amount of air
9 they're breathing in and the potential for
10 absorption.

11 Okay. So now I am going to change gears a
12 little bit and switch to the biomonitoring
13 data, and I have to tell you I am going to
14 focus on one compound, pyrene. Pyrene is one
15 of those PAHs that was in the first slide.
16 It's an important component of PAHs. It -- of
17 -- and it's representative of the four and
18 five ring carcinogenic PAHs, okay?

19 So, of all of those type of compounds, pyrene
20 is the most abundant. So it's oftentimes the
21 easiest measured, and we do have a biological
22 exposure indices for 1-hydroxypyrene, the
23 major metabolite of pyrene, which is an ACGIH
24 BEI. That was developed in -- I'm not sure it
25 was in place in 2001. It may have been.

1 We'll have to go back and check that when we
2 think of it.

3 But biomonitoring can account for differences
4 in absorption, distribution, and metabolism
5 and elimination if it's done correctly. It
6 can take into account both the skin and
7 inhalation exposures and one very important
8 thing with biological monitoring is that
9 exposure can be reconstructed.

10 If you know the material that you are exposed
11 to and you know the half-life of that material
12 in the body and you know the relative time
13 between when the sample was taken and when the
14 exposure occurred, you can reconstruct the
15 exposure based upon the half-life.

16 On the other hand, it is a method that is
17 easily misused, if not in terms of
18 interpretation, if you don't know exactly what
19 you're doing, so.

20 Let's look -- and this is an example of a
21 biological monitoring on a model system. This
22 has nothing to do with the Trade Center. This
23 is just a model that I made up. So you see if
24 you have exposure on Monday morning and the
25 exposure during the day on Monday equals to a

1 hundred, and the half-life in the material in
2 the body is 24 hours, then the material -- you
3 will increase the amount in the body, and then
4 in the 16 hours the person is off until the
5 next shift on Tuesday, that level will
6 decrease by a fraction based upon the half-
7 life.

8 So you can see right that you get a -- with
9 each additional day, you get an increase, but
10 it's not a doubling. So you don't get 200 on
11 Tuesday; you don't get 300 on Wednesday and so
12 forth. And then the other thing to notice is
13 that because of the half-life -- and what is
14 half-life?

15 Half-life is -- most of you probably know --
16 is the length of time a material resides in
17 the body. Most of the materials that are
18 absorbed by humans as xenobiotics are
19 eliminated. And they are eliminated fairly
20 rapidly because the body doesn't want to keep
21 these things if they do nothing for it. I
22 mean, some materials have long half-lives;
23 cadmium has a 30-year half-life. Lead has
24 about an eight-to-ten year half-life in the
25 bone. But these materials tend to be

1 eliminated fairly quickly and with fairly
2 well-defined half-lives.

3 Notice what happens after work on Friday. So
4 after work on Friday, the level in the body
5 goes way down before Monday morning, and
6 that's because there are several half-lives
7 involved here, okay. So when would be the
8 best time to sample for this material,
9 something with a 24-hour half-life?

10 Now you wouldn't want to sample on Monday
11 because the body hasn't reached steady state
12 yet. Oh, and by the way, this continues every
13 week. It doesn't get much higher. It never
14 gets above 200 for this compound as long as
15 that dose is the same.

16 When would you want to sample? Well, you
17 don't want to sample here. You really want to
18 wait until the end of the week. Sample in
19 here and you'll have less variability, and
20 you'll capture the exposure because that's
21 when the exposure reaches its peak.

22 You wouldn't want to sample down here at this
23 time because that would -- without knowing
24 when the peak occurred -- because that would
25 underestimate exposure dramatically. So let's

1 look at the data. These are the 1-
2 hydroxypyrene data from Edelman et al, and
3 this is one table I looked at, and I'm only
4 giving the 1-HP data. And I've changed the
5 numerals that have been used, and in that I
6 use micrograms per liter and I'll tell you why
7 momentarily.

8 They use nanograms per liter. Micrograms give
9 smaller numbers, fractional numbers, but it's
10 important because the BEI is set at one.

11 Okay, so all exposed workers at the site when
12 they were sampled on October 1st, 2nd, or 3rd
13 had a level of 0.092 micrograms per liter.

14 The controls had a level of 0.062 micrograms
15 per liter, and that seems like a small
16 difference, but it could be a significant
17 difference and it was in fact significant. It
18 was significantly higher.

19 If the firefighters were at the collapse on
20 day one, then their average was about .11. If
21 they were -- if they didn't come at the
22 collapse, but came after the collapse on day
23 one and two and started working, then it was
24 slightly, slightly higher, so maybe if you
25 could say the real fires that were happening

1 at ground level didn't happen until here, at
2 least in the majority of the -- after the
3 collapse. That's when all hell broke loose.
4 There was a subgroup that was studied which
5 was called the Special Ops Command, and they
6 were considered to be the highest exposed, and
7 indeed, they had the highest average level.
8 Their level was .159.

9 Okay, now the reason when I looked at these
10 data initially I thought that well, you know,
11 you can see there's a significant difference
12 here but it's not a big deal, was because the
13 standard that occupational exposures are based
14 on, the level is 1.0, okay?

15 So the occupational standard is much lower,
16 but it specifies an end of shift, end of
17 workweek sample and as I found out by reading
18 the paper hard, one, they did not capture the
19 peak. Samples were collected 20 some days
20 after the exposure, which would be -- and also
21 they reported no variances and other people
22 can maybe reinforce this, but when we were
23 worried about people who have exposure, it's
24 the outliers that are really important, and
25 the outliers weren't given in the paper.

1 Four percent were said to be in the upper five
2 percent of the NHANES values, but I wonder how
3 many of the controls were in the same upper
4 five percent. It wasn't represented. Because
5 then there's no comparison there. But there
6 was no variation given. There was no standard
7 deviations, no ranges that were given in the
8 data, and no exposure time was indicated or no
9 sampling time was indicated. They did not
10 indicate whether they sampled at the end of
11 the shift, at the beginning of the shift or
12 when they sampled at all. It's just unknown,
13 and that really threw me, okay?

14 So we have a situation where the exposure may
15 have occurred many days before and also -- and
16 so you would expect them to be relatively low
17 relative to the decrease in exposure that one
18 might see with that decrease in the PAHs that
19 were reported.

20 Going back to the -- if I may, this slide.

21 So, regardless of what the true levels were if
22 these were just area samples, you can see that
23 the shapes of the curve are similar. So one
24 may anticipate that if there was a higher
25 level inside of Ground Zero, then it would

1 follow a similar shape, so the levels that --
2 this is when the -- the highest level would
3 have been reported here. The first samples
4 weren't taken to here, out 25 days, and you
5 can see what the shape of the curve looks like
6 in terms of the exposure. It's already
7 winding down at least.

8 Now how can we -- can we do anything with this
9 data and -- okay. So the sampling time wasn't
10 given. Firefighters -- and this is from my
11 own experience that firefighters haven't -- in
12 the studies that we've done in Cincinnati, the
13 firefighters have a higher level after a fire
14 than before, but generally they are not in the
15 really high exposed level and I'll give you an
16 idea of what that means here in a moment.

17 And then the question becomes are -- could
18 absorption from the lung be complete? What
19 about the large particle masses and the fact
20 that PAHs might not be absorbed rapidly, and
21 I'll show you some data on that in a moment.

22 So first things first. This is what happens
23 in a workplace in an aluminum plant, and I
24 showed you what those look like. In aluminum
25 plant workers, and their exposure to 1-

1 hydroxypyrene. These samples were taken pre-
2 shift, so there was a baseline sample taken
3 every morning and an after work shift, and you
4 can see that their exposure follows the model
5 for a 24 -- very similar to what I reported
6 earlier.

7 But look at the magnitude of their exposures.
8 By the end of the workweek, these levels are
9 greater than 10 micrograms per liter -- per
10 liter of urine, which is 10 times the
11 standard. But notice that every day before
12 the shift, they drop down considerably, so
13 that if this is the peak -- and what this
14 shows is that like in many workplaces,
15 aluminum reduction workers don't produce as
16 much on Friday as, you know, it's Friday.
17 But you can see that after Thursday's peak,
18 that there is a significant drop in the 16
19 hours between the next day. So if you didn't
20 sample, if you sampled in the morning, you
21 would see a much lower sample by design, much
22 lower level by design. And these are data
23 that were developed by the BEI committee in
24 running up, in developing the BEI for 1-
25 hydroxypyrene.

1 And what they show -- it looks complicated,
2 but what it shows is how exposures could be
3 the sum of all of the different compartments
4 for these things. It's known that PAHs have
5 three compartments in the body: the blood,
6 which is cleared very rapidly with a half-life
7 of five hours; the lean tissues, which are
8 cleared within 24 hours; and then the --
9 probably the adipose tissues which are cleared
10 very slowly, just every -- the half-life is 23
11 days approximately.

12 And so what you see is that with every
13 exposure, the major impact on the urinary
14 levels shown in black is the sum of the three
15 of them, but it's largely dependent on the
16 lean compartment and the -- and what was in
17 the blood, and then that rapidly disappears
18 causing a drop in the urinary levels.

19 This was an example I found extremely
20 illuminating for this discussion. This was a
21 group of people, patients in this case, who go
22 to the Mayo Clinic for what's called the
23 Goeckerman treatment where they have psoriasis,
24 and their skin is painted with as much as 70
25 percent of the total body volume of -- their

1 skin is painted with coal tar in the treatment
2 of psoriasis. It apparently works.

3 And what I'd like to focus on -- the slide is
4 more complicated than it needs to be. I'd
5 like you to look at the -- the values here for
6 1-hydroxypyrene. So these are the baseline
7 values in this group of people. After one
8 treatment, that baseline jumps up to 170,
9 okay? Now this is applying it on the skin.
10 After five treatments, because they're given
11 eight hours a day of this treatment, five days
12 a week, and then it's stopped. After five
13 treatments, it goes up to 270, approximately,
14 but after one week of no treatment, this is
15 the level. And it goes down -- remember
16 there's a break here between 10 and 100 -- and
17 it goes down between 275 and down to less than
18 4 within a week.

19 If you calculate that, that means that the
20 half-life for this is about 24 hours, which is
21 very consistent for a group of people who
22 haven't been exposed chronically. Their
23 exposure was just five times. So it drops
24 very rapidly with an apparent half-life of
25 about 24 hours.

1 Why this is important is that if the half-life
2 was indeed 24 hours, one could back calculate
3 from the levels that are given to the levels
4 that may have been at the peak on 9/11, 9/12
5 at Ground Zero.

6 What this slide shows is the data from Gerde
7 et al, who looked at the impact on particle
8 size. PAHs were absorbed onto particles and
9 then they -- and then they modeled it into the
10 lungs based on -- and then actually did actual
11 measurements in the lungs, and what they saw
12 was the smaller the particle that the PAH was
13 held on to -- so these are particles with PAHs
14 on them -- when they were deposited in the
15 lung, a very small particle had a very short
16 half-life.

17 So if it was .1 micron, the half-life is
18 approximately less than a minute, probably 30
19 seconds; but if it was a very large particle,
20 the half-life could be more -- much more
21 extensive. So we're talking on the orders of
22 a month or greater if it was 1000 microns.

23 Now how might a particle get to be 1000
24 microns in the lung? Imagine that -- and what
25 we used to see in tobacco smokers was that

1 you'd get these agglomerations of tars at the
2 bronchial -- where the bronchia would split
3 and tars would accumulate, and that makes the
4 particle much larger and makes absorption from
5 it much smaller.

6 So the idea is that an exposure even one time
7 can result in a very prolonged exposure based
8 upon the fact that it comes off a larger
9 particle much slower.

10 Then there's the part of how with the amount
11 of deposition, and I'm not going to go too
12 long in this, but what it really shows is that
13 if you breathe regularly, you -- regardless of
14 the particle size, this is the fraction that's
15 collected and deposited in various areas. But
16 if you breathe a lot faster with a much higher
17 tidal volume, breathing in deeper, then you're
18 much more effective at collecting particles.

19 So people who are working harder not only
20 breathe in more air, but they also deposit
21 much more readily.

22 So PAHs do absorb on particles. Soot,
23 particularly, so on diesel exhaust and those
24 types of things, they -- because of their
25 lipophilicity, they are very much attracted to

1 those soots. But they are also attracted to
2 concrete particles, and that's been shown in
3 the literature, to a lesser extent, but still,
4 they're absorbed onto the particles and then
5 deposited and held in the lungs.

6 The particles may accumulate in the lung and
7 slow their absorption into the body, and
8 particles may be coughed up, expectorated,
9 spit, or swallowed, but this, in fact, seems
10 to be more of a detoxification pathway than an
11 exposure pathway for a complicated reason
12 dealing with the liver first pass. Okay, you
13 know what I mean, but...

14 On the other hand, PAHs have known to interact
15 with other exposures. PCBs and dioxin were
16 found on the site. In fact, the highest
17 ambient level of dioxin ever measured was
18 measured in the world after 9/11. Dioxin is
19 known to be used as an enhancer of the
20 carcinogenicity of some PAHs, so if animals
21 are treated with dioxin, they are more likely
22 to get tumors than if they're not treated with
23 dioxin and given the carcinogen.

24 Silica is something that we haven't mentioned
25 too much, but PAHs are known to enhance the

1 carcinogenicity of silica exposure. And in
2 this case, when I'm talking PAHs, I'm really
3 talking smoke. The interaction seems to be
4 additive or additive plus, and then unlike
5 what John mentioned, the data that I looked at
6 saw that PAHs, again, smoking, enhanced the
7 carcinogenicity of asbestos, but at least the
8 studies that I -- the consensus was that it
9 was multiplicative but I would certainly --
10 he's much more experienced in this than I am.
11 So the conclusions that I would make are that
12 exposures to workers to PAHs within the Ground
13 Zero site was almost certainly higher and
14 maybe substantially so than was indicated by
15 the majority of exposure studies. A fuller
16 report of the biological monitoring data is
17 needed to predict what exposures may have been
18 during the early periods after 9/11 and who
19 may have been at the highest exposures.
20 The people who are the outliers are the key.
21 If the people who had the highest levels of 1-
22 hydroxypyrene are the ones who later -- they
23 have the highest dose, and they may be the
24 ones who are at the highest risk, and
25 understanding who, not who the outliers are

1 from our point of view, but what the range of
2 the outliers were and then moving that back is
3 an extremely important thing, at least in my
4 mind.

5 And if the effective half-life is 24 hours,
6 then the 1-hydroxypyrene levels on 9/12 could
7 have been well above the BEI assuming that
8 there was no exposure, assuming that there was
9 no exposure. Now, that's not the case. There
10 was exposure afterwards.

11 The best thing to do would be to model that
12 exposure, and the half-life would be -- with
13 the curves that were used in the exposure
14 studies. You'd have to integrate those
15 together. I didn't have the time to do that,
16 and I -- yeah. It's something that one could
17 do, though. Thank you.

18 MS. FLYNN: Thank you, Glenn. A quick
19 question. What would the exposure metrics be
20 for a 10-year-old child?

21 DR. TALASKA: No idea. I'm sorry, I shut it
22 off, and I killed it. I've got it. I have no
23 idea.

24 MS. FLYNN: Because in general, as I
25 understand it, and maybe Leo could comment on

1 this, but children actually take in more air
2 than adults, so I wonder --

3 DR. TALASKA: Well, again, and you do have to
4 realize that at the fence line, they were
5 measuring those exposures and the exposures
6 were tending to rise. I can't tell you, but
7 kids weren't inside of Ground Zero, okay, so I
8 don't know what the exposure would be because
9 the data are so -- but kids tend to breathe
10 more. They have larger surface area relative
11 to their body, so they do tend to sometimes
12 take in more materials. They do eat things.

13 MS. FLYNN: Kids were not inside of Ground
14 Zero, but, we actually, you know, do have
15 available -- I'd have to find them on the
16 site, the High School Parents Association
17 website, but information that show that on
18 days when debris was being dumped on the
19 hazardous debris barge outside of the
20 Stuyvesant High School ventilation system, the
21 particulate concentrations were comparable to
22 Ground Zero.

23 So, I mean, there were lots of -- there was
24 just tremendous potential for different kinds
25 of exposures that have not been captured in

1 the data, so we just -- this is something that
2 -- I know I sound like a broken record, but I
3 think it's really, really important to keep in
4 mind number one, number two. Children were
5 caught in the dust cloud in the initial
6 collapse cloud, so I don't know if Leo if you
7 want to add anything.

8 DR. TALASKA: I didn't look at that. I'll be
9 honest. I was focusing -- there was more than
10 enough here to cause me to -- so I really
11 didn't look at that in a really hard way.

12 MS. FLYNN: Can I just make a plea on behalf
13 of the stakeholder members of this panel? We
14 actually -- we're not experts and we obviously
15 defer to the scientists here, but we're equal
16 members of the panel and we know a lot of
17 things because we've been basically engaged
18 with, you know, the facts on the ground from
19 the very beginning.

20 So if it's possible for us to have in advance
21 the drafts of your presentations -- I'm sorry
22 I keep popping my keys -- the drafts of your
23 presentations, that would be tremendously
24 helpful. I know that Susan Sidel provided
25 extremely valuable information to -- to

1 Virginia Weaver, and we want -- we didn't want
2 to load you guys up, because we know that, you
3 know, you're like, you're trying to condense a
4 tremendous amount of material, but there were
5 times when we actually can bring a useful
6 perspective and we really appreciate that
7 opportunity.

8 MS. HUGHES: Also, it seemed like most of the
9 sampling was done at street level, and if you
10 look at the topography downtown, it's
11 surrounded by very large skyscrapers. So if
12 the plume actually expands would the results
13 of the testing might be different higher up?
14 You have families living in these high rises
15 in very close proximity, so I just wanted to
16 mention that as an exposure route.

17 And the second thing is, it wasn't as though
18 the only fire was where the two towers were.
19 It spread, and you had gas lines feeding --
20 pardon me -- but there was gas lines feeding
21 the World Trade Center site. So there is
22 exposure within the area, and it went on and
23 on and on, so I just wanted to put that in for
24 the record.

25 DR. WARD: I suggest -- a suggestion, we are

1 running late, and maybe we'll take one more
2 comment and then we'll have a 10-minute break
3 and then resume, because we do have a fixed
4 time when we need to start the public
5 comments.

6 DR. MARKOWITZ: So, John, John Dement made a
7 point on discussing asbestos as there is no
8 known safe threshold. So the question, since
9 you frame the exposures among the firefighters
10 around the biological exposure index, what's
11 the relationship between the BEI and cancer
12 risk for PAHs?

13 DR. TALASKA: It's not really known. The BEI
14 is based upon specifically the level that is
15 associated with occupational exposure if you -
16 - and not with environmental exposure. There
17 wasn't sufficient data to be able to say that
18 there was any level of -- that was related to
19 disease yet.

20 There weren't simply enough data there. There
21 are data that shows at that level since then -
22 - we've put out -- we've done studies showing
23 that at the level of the BEI of one microgram
24 per liter, there's an increase in PAH, but we
25 don't know what it is relative to cancer as of

1 yet.

2 There aren't sufficient data, but -- so that
3 the level was set just so that it would rule
4 out things like tobacco smoking because you
5 can't get -- smokers don't have levels that
6 are that high, as high as you want. Does that
7 answer?

8 DR. WARD: Okay, we'll take a 10-minute break.

9 (Recess taken from 2:52 p.m. until 3:12 p.m.)

10 DR. WARD: Let's start. I think everyone's,
11 virtually everyone's back at the table and we'll
12 start with the presentation by Bill Rom.

13 **PARTICULATES AND WTC**

14 DR. ROM: Thank you, Elizabeth. Does Paul have
15 some slides? My task is to talk for five minutes
16 about particles, particulates or particulate
17 matter. My job is to talk about exposure
18 assessment, what were the exposures; second, how
19 bad are these particles, are they really toxic or
20 are they not toxics; and third, what is the
21 evidence for these particles in humans, did they
22 get exposed and how much; and lastly, for gravy,
23 are these particles going to cause cancer, since
24 that's the question we have to address soon.
25 On this slide you see the particles on the left and

1 then you see the fires on the right. The point I
2 would like to make is that there were two kinds of
3 exposures here, but I don't want to make that point
4 so much as to say that they overlap. This was a
5 fire that was extremely hot, that burned the
6 particles, and we have a particulate exposure that
7 really has never been seen before. This is unique.
8 This is a disaster medicine and these particles
9 really can't be classified basically like coming
10 from the mine or source 'cause they've been
11 altered.

12 Next slide. So this is a grab sample of the dust
13 particles on the right. This is WTC dust but a
14 third of that dust comes from wallboard. So all
15 this stuff that we're seeing right there. So
16 that's gypsum, and gypsum is calcium sulfate. It's
17 not -- it's what we always call with NIOSH,
18 nuisance dust. We chuckle about that 'cause we
19 wonder what it is. Calcium sulfate is not known to
20 be very toxic; it's mixed in with calcite. Calcite
21 has calcium carbonate and calcium carbonate is not
22 very toxic, but it forms little crystals and when
23 you see it in tissue, can actually be birefringent,
24 and that's important to remember in regard to
25 silica.

1 Third, there is some cement dust mixed in here and
2 the cement dust is calcium hydroxide. And that is
3 a basic salt and it's alkaline, so we know the pH
4 of this World Trade Center dust was around 11 so
5 it's alkaline and it's irritating. It's irritating
6 to the mucus membranes, to your eyes, to your
7 mouth, to your throat, makes you cough. So is that
8 really something that's going to cause lung disease
9 and cancer?

10 I had the good fortune of being funded by NIOSH to
11 study trona miners, and trona miners were exposed
12 to a sodium sesquicarbonate that we use for the New
13 York Times and Coke bottles and things like that.
14 And the trona mines are in Wyoming, so I had to go
15 to Cheyenne and have a personal interview and get a
16 medical license, and then spend a couple weeks in
17 Rock Springs and Green River with cowboys, and they
18 would mine trona.

19 So we studied 230 trona miners and we looked at
20 shift studies to see if they would have a drop in
21 lung function over shift and any alterations in
22 their breathing, and it was really a negative
23 study. So pure trona, sodium sesquicarbonate, is a
24 rather benign dust.

25 But they all complained of skin itching and

1 dermatitis and irritation, and we got a second
2 paper on just trona dermatitis. So that shows you
3 that alkaline dust can irritate the mucus
4 membranes. So in its pure form these dusts are
5 rather benign.

6 But then you also notice on the left of this slide
7 that a lot of this dust was respirable, less than
8 2.5 microns, that's not mm, it's microns, so
9 there's a lot of respirable dust that gets down
10 into the lungs.

11 Last week Dr. Jim Ha was visiting us at Bellevue,
12 and we spent an hour looking at eight lungs that
13 were from open lung biopsies of World Trade Center
14 dust exposed people, and we looked for silica and
15 we really didn't see birefringent particles sharp
16 and bright like silica, so I'm going to dismiss
17 silica as really being a critically important
18 particulate exposure to the workers. And I'll
19 point that out by looking at the next slide.

20 So we've documented an exposure and now I want to
21 go on to the toxicity of these particles. So we
22 had a firefighter who came within the second week
23 of 9/11 to Bellevue who was critically short of
24 breath and ended up in the medical ICU, and he had
25 bilateral infiltrates and effusions, and we didn't

1 know what he had so he was treated with antibiotics
2 and steroids, and was getting better. But since
3 I'm a physician-scientist and I'm the boss, I like
4 to yell at my faculty, I said, you need to get him
5 consented and do a bronchoscopy, you know, lavage
6 and make a diagnosis.

7 So fortunately he agreed to the consent and we were
8 able to get some cells. And he had all those red
9 cells on the right, that's acute eosinophilic
10 pneumonia. So he had a very unusual disease that
11 may be related to dust exposure. The important
12 thing is we got those cells and you can see they're
13 pretty clean. They don't have smokers' particles
14 in them, so we sent these cells on the next slide
15 to Victor Roggli down at Duke to analyze them for
16 particles. And we said, this is a firefighter
17 exposed for two weeks in the Pile, and this is the
18 first lavage, and these are cells from his lung and
19 we want to know what particles are down there.

20 So first of all, he showed us a fiber, and that's
21 an amosite fiber on the left because he did an
22 x-ray dispersive analysis for elements and found
23 iron as well as magnesium and silica, and pointed
24 out that that's an eight-micron-long fiber.

25 The important thing is it's not coated. It's an

1 uncoated fiber which means it's freshly inhaled,
2 which is very unusual. You never see that in
3 asbestos workers unless they're from the mines in
4 Quebec.

5 The middle particle I want to point out to you, is
6 what I think is a really toxic WTC particle 'cause
7 that is something that looks like from outer space.
8 I called it fly ash particle 'cause it reminds me
9 of a clinker coming out of a coal fire. But I
10 think that's a burned particle. And in your packet
11 there's an analysis of particles from the Deutsche
12 Bank building, and the analysis shows a lot of
13 these particles are coated with other substances
14 from the fire, and that probably enhances the
15 toxicity of these particles, so that's a burned
16 particle.

17 On the right is what we think is fibrous glass, and
18 you can see it's not parallel on its sides. It's
19 probably been exposed to 100 degrees temperature so
20 it's been partially burned.

21 The fourth thing I want you to look at is on the
22 bottom. There's 305 commercial asbestos fibers per
23 ten to the million macrophages. So how much were
24 these people exposed to? So in my tenure at the
25 NIH, I lavaged about 500 coal miners and asbestos

1 workers and silica exposed workers, and I had to do
2 some normal volunteers. So I had eight normal
3 volunteers and they had a mean of 30 asbestos
4 fibers per million macrophages. So this
5 firefighter has about ten times the normal number
6 of fibers in his macrophages. And the asbestos
7 insulators I would lavage would have about a
8 thousand. So he's, you know, just after a couple
9 weeks, he's up to a third of the way to what an
10 insulator has in his lung.

11 Now, I would say that breathing the air with your
12 nose and your lungs is probably a better
13 measurement than the samples that EPA took, and we
14 couldn't find any fibers in their samples. So this
15 guy was on the Pile and trying to rescue that --
16 this whatever could be done to save others.

17 Next slide. So this is what chrysotile asbestos
18 looks like, and the reason there was an amosite
19 particle there, is that in New York, when we put
20 chrysotile asbestos in the sprays and on the steel
21 girders, we always threw in about five percent
22 amosite. Reasons, I don't know why but they always
23 did that so that's why you find a mixture.

24 Next slide. So this is from the asbestos
25 insulators and the kind of fibers you normally

1 find. That fiber has a coated iron and protein
2 surface and that's what those beads look like. So
3 this is a fiber that's been sitting in an insulator
4 for 20 or 30 or 40 years. And you see the body
5 tries to protect itself by walling off the fiber.
6 And the other cells are macrophages, and this is a
7 nonsmoking asbestos insulator, and there's no other
8 particles in there. So he's a clean asbestos
9 insulator from being nonsmoking, at least. Not
10 clean in terms of fibers.

11 Next slide. So Dr. Selikoff taught a number of us
12 in this room about asbestos insulators, and his
13 very famous study about all of the North American
14 insulators showed a five-fold increase of lung
15 cancer and almost 10 percent had mesothelioma.

16 Next slide. And when I was at the NIH I would
17 spend weekends recruiting patients for a lavage,
18 and I would sit with Jim Keogh at the Baltimore
19 City Hospital recruiting in study subjects, and he
20 had one of his patients from Sparrows Point Steel
21 Mill who had silicosis, those are the nodules on
22 the right, and he also had mesothelioma with the
23 left, if you reverse looking at this patient, with
24 a big pleural effusion. So mesothelioma is the
25 other disease along with lung cancer that you get

1 from asbestos. How much asbestos causes
2 mesothelioma, I remember when I was working for
3 Dr. Selikoff, he had me interview a 55-year-old man
4 with mesothelioma, and he worked in a flower shop
5 in Brooklyn, and I couldn't figure out any reason
6 he got mesothelioma from flowers. And I remember
7 that in Tyler, Texas, the flowers came in gunny
8 sacks and maybe the gunny sacks were used for
9 asbestos. I asked him about gunny sacks, he said I
10 don't know. I never saw gunny sacks. Then I asked
11 him if he worked in the shipyard, and he had worked
12 in the Brooklyn Navy yard for one summer in 1942 as
13 a helper, and had two and a half months of shipyard
14 exposure. So very minimal exposures can cause this
15 disorder.

16 Next. The marker for asbestos are pleural plaques,
17 the blue and purple around this lung are pleural
18 thickenings.

19 Next slide. And if you have those, Hillerdal in
20 Sweden showed that if you have pleural plaques, you
21 have a slightly increased risk for lung cancer and
22 an increased risk for mesothelioma, so this is a
23 marker of your asbestos exposure.

24 Next slide. And importantly, Dr. Selikoff would
25 take us to Paterson, New Jersey, where there was an

1 asbestos factory, making fire hoses for New York,
2 and he followed a hundred men who worked for just
3 two months, from 41 to 45 in this factory, and
4 followed them to the end of the 1970s. And on the
5 right you can see with the dotted line that 25
6 years the lung cancer observed rate increased over
7 the expected, so just for two months of exposure 30
8 years earlier, you have an increased risk for lung
9 cancer.

10 The project that I was involved in was doing lung
11 function on the wives of these workers. And I did
12 about 300 spirometries showing that they had a
13 reduction in their spirometry from doing the work
14 clothes washing of their husbands and hugging them
15 when they came home from work from Paterson's
16 factory. And among those wives, four of them ended
17 up getting mesothelioma from that exposure.

18 Next slide. So Dr. Ward wanted me to go over
19 particles and lung cancer, so the small burn
20 particles that we have from diesel exhaust have
21 been studied in the American Cancer Society cohort.
22 The American Cancer Society enrolled over a million
23 adults in 1982 about the risk for cancer. But
24 these people lived in metropolitan areas throughout
25 the U.S. that had EPA-collected data on particulate

1 matter of 2.5 microns in size. So almost half of
2 this cohort had data on particulate exposure
3 through the end of 1998 from 1982.

4 So in the next slide on the left, you can see the
5 lung cancer mortality. On panel A is
6 cardiopulmonary mortality; panel B on the lower is
7 lung cancer mortality. The three circles on the
8 far left are above the line of 1.0 so all three
9 dots are statistically significant over time for an
10 increased lung cancer mortality of approximately
11 8 percent from PM(2.5) exposure, which is the burn
12 particles from diesel exhaust.

13 Next slide. And these are what the particles from
14 diesel exhaust look like in macrophages from the
15 lung. This is a collection from sputum in children
16 in England. And these macrophages were looked at
17 under a light microscope and you see the black
18 particles, particularly in D and E, that are very
19 tiny, less than 2.5 microns.

20 The next slide, we'll skip and go to the slide
21 after it. These are from families, next slide,
22 that did not have any smokers in the household and
23 they were on at least a second level, so they were
24 a little bit away from the street level. And on
25 the slide on the upper left you'll see a declining

1 FEV-1 in those children as they had increased
2 numbers of those particles in their macrophages.

3 Next slide. So these diesel particles cause
4 adverse health effects.

5 And lastly is cancer. So cancer in the lung starts
6 off as abnormal proliferation and survival of
7 injured cells in the respiratory epithelium
8 associated with genetic defects, whether they are
9 specific genes that are up-regulated, down-
10 regulated, insertions, deletions, mutations,
11 amplifications and so on, that you end up getting a
12 clone of cancerous cells.

13 Next slide. And the last point I'll make is that
14 there are now ways to diagnose these cancers with a
15 blood test. And you can now target proteins in the
16 blood to diagnose these cancers. On the top in the
17 white are little aptamers, that are nucleic acids
18 designed to pick out a protein in the blood, and
19 you can make more than a thousand of those aptamers
20 to pick up specific proteins in the blood.

21 And next slide. This assay has been looked at in
22 1300 lung cancer patients and matched controls, and
23 you can see that a panel of about 13 biomarkers can
24 very accurately pick out the lung cancers with area
25 under the curve of .9. So in looking forward at

1 lung cancer and mesothelioma, there are tests at
2 the early and past research level to identify these
3 people both at risk and of getting the disease.
4 And this test is about to be commercialized for
5 mesothelioma as the first disease to look at.
6 I think that's it.

7 DR. WARD: Questions or comments for Dr. Rom?

8 (no response)

9 **METALS, VOCs and WTC**

10 Okay, is Virginia on the line?

11 DR. WEAVER (via telephone): Yes, I am.

12 DR. WARD: Are we ready to...

13 DR. WEAVER: I am ready. Can you guys hear me if I
14 stay on speaker phone?

15 DR. WARD: Paul just cautioned me that we only have
16 14 minutes before the -- before the public
17 presentation -- public comment period. And so why
18 don't we get started and see if we can wrap up your
19 presentation in that time frame and then if
20 necessary, can you come back and we can have
21 questions after the public comment period?

22 DR. WEAVER: Yes.

23 DR. WARD: Okay, great.

24 DR. WEAVER: You have my slides up?

25 DR. WARD: Yes, we've got the first one up.

1 DR. WEAVER: So after the title slide, moving to
2 the second slide, I wanted to simply give you some
3 of the thoughts that were going through my mind as
4 I was looking at data related to volatile organic
5 chemicals and metals. And one issue in my mind was
6 the shortest exposure duration that results in a
7 measurable increased risk for cancer, and I've been
8 very happy to hear discussions about increased risk
9 in very short time period. I was not aware. I'm
10 not a cancer expert, and I was not aware about that
11 data, and that's very helpful to us in thinking
12 about risk from exposures that are of -- that occur
13 only when you're actively exposed, which would be
14 the volatile organic chemicals.

15 The other point that I was thinking about as I
16 prepared these slides are that we are now learning
17 that a steeper exposure rate may result in greater
18 risk, so for the same overall accumulative dose, if
19 you get the exposure faster, the risk may in fact
20 be greater. And so what that means is that the
21 exposure construct for cancer outcome differs from
22 that that's been used in World Trade Center
23 research for pulmonary outcome, so rather than
24 looking at where you were at the time of the
25 collapse and shortly thereafter, we have to think

1 about burning tile, diesel exhaust and carcinogens
2 in dust.

3 So on the next slide I had simply shown an example
4 of one type of exposure characterization and I know
5 Liz has already showed this type so I'm going to
6 move right on to the next slide on key concepts and
7 questions.

8 We've already heard that cancer of course varies by
9 time since exposure onset, and so it is the
10 nonsolid tumors that are the ones we could be
11 seeing, even at this point, from World Trade Center
12 exposures but specifically the leukemias. And then
13 a point that I think others have already made so
14 far is that we have very little data about chemical
15 mixtures overall, particularly in the World Trade
16 Center yet. This is a common exposure scenario
17 overall and of course clearly at World Trade
18 Center.

19 The next slide I simply wanted to show the group 1
20 and 2A IARC carcinogens that are in the volatile
21 organic chemical category. I took this from
22 NIOSH's summary. I want to point your attention to
23 benzene, which has been classically linked to what
24 we used to call acute myelogenous leukemia but we
25 now call acute nonlymphocytic leukemia as our

1 ability to analyze these types of cancers has
2 improved.

3 I also want to point out that there is limited
4 evidence that benzene causes acute lymphocytic
5 leukemia, chronic lymphocytic leukemia and
6 importantly multiple myeloma. That is from IARC
7 and it's also supported by a meta-analysis
8 published in EHP in 2008, again, supporting that.
9 Other VOCs that were of concern from World Trade
10 Center would include 1, 3-butadiene, which is a
11 combustion product like benzene, from the Pile and
12 also from diesel exhaust. Again, this has been
13 linked to leukemia and also non-Hodgkin lymphoma,
14 formaldehyde, nasopharyngeal cancer, and there's
15 increasing evidence that formaldehyde is linked to
16 leukemia as well. That's considered strong but not
17 sufficient evidence based on the NIOSH summary and
18 vinyl chloride. And then I've listed some of the
19 2A, which are -- Group 1 of course is known human
20 carcinogens, Group 2A is, I think the
21 categorization is probable, and it's based on
22 adequate animal data but inadequate or limited
23 human data.

24 So in the next slide, the important aspects about
25 exposure to VOCs is that they're common in

1 combustion products. I think about this a lot in
2 the work I do for the firefighters union. So you'd
3 think about this from working on the Pile, from the
4 smoke and exhaust from that, and also diesel
5 exhaust.

6 In general VOCs, as the name implies, are not
7 persistent in the environment and they do not
8 accumulate in the body so the exposure duration
9 would have been while you were actively working on
10 the Pile. But also importantly, these exposures
11 are associated with some of the shortest latency
12 cancers, ones that we could be seeing.

13 Next slide. As far as I can tell, and I'm no
14 expert on World Trade Center exposures, there are
15 very limited data on VOC measurements. There were
16 grab samples that were taken on the Pile to try and
17 determine if it was safe for rescue workers to
18 enter. So Lorber et al noted that when samples
19 showed, quote, extremely high concentrations of
20 VOCs, end quote, entry was prohibited. I don't
21 have levels about exactly how high those were.

22 Lorber notes that for a number of the VOCs found
23 elevated levels outside of Ground Zero but still
24 within restricted zones, and when they used 24-hour
25 samples, which should give a little bit better

1 measure. You know, generally in a work place we
2 measure eight-hour samples. When they compared
3 grab samples over four minutes to 24-hour samples,
4 they found that levels were much, much lower for a
5 number of the VOCs of concern, including ones from
6 butadiene. However, that was not the case for
7 benzene. The benzene monitoring showed many more
8 grab samples that were higher and 24-hour samples
9 that, rather than being a thousand times lower,
10 were about ten times lower.

11 I'm not sure if I said next slide but I have a
12 separate slide on benzene monitoring. And on that
13 slide I included the samples for benzene in 24-hour
14 measurements that were above the detection limit,
15 and so apparently there were only fourteen 24-hour
16 samples that were done for benzene, which doesn't
17 seem like many. Six were above the detection limit
18 and of those, a few were fairly close to the Agency
19 for Toxic Substances and Disease Registry
20 intermediate minimal risk level, which would apply
21 for folks who were working for more than a month,
22 more than 14 days up to a year.

23 In the conclusion in the Lorber article, which as
24 the data suggests in exposures to benzene at levels
25 that approach the intermediate MRL were not likely

1 to have lasted longer than 45 days.

2 There's a few samples from truck drivers, done by
3 my colleagues at Hopkins, that were not
4 extraordinarily high either. You know, in the low
5 parts per billion compared to workers are allowed
6 to be exposed a thousand parts per billion.

7 And I was going to make the point with the text
8 below that the monitoring levels seem inconsistent
9 with the descriptions and pictures of the site, but
10 I think others have already made that point more
11 eloquently before me. There is an inconsistency
12 between monitoring and what was visualized.

13 So in thinking about the potential implications of
14 VOC exposures, in my mind it would be workers who
15 were on the Pile would be at most risk, and
16 obviously the longer they worked on the Pile, the
17 more risks they would incur.

18 I was thinking about how much time you would need
19 to work there in order to have increased or
20 measurable increased risk, and with the
21 understanding that probably the exposures were
22 much, much higher than any of the monitoring data
23 that we have. And so I guess it would be a matter
24 of thinking about individuals near and on the Pile
25 and the length of time that they worked in those

1 capacities and that would be how we would consider
2 risk relating to VOCs as an important consideration
3 because this exposure that could be resulting in
4 cancers early on.

5 And then I'm going to shift gears and talk about
6 metals so that's the next slide. There are a
7 number of metals that have been associated with
8 carcinogenicity in a variety of different organs.
9 I've listed those for you here, again, from the
10 NIOSH summary document.

11 On the next slide, I want to step back quickly and
12 thank Susan Sidel for helping me come up to speed
13 over the course of the weekend on World Trade
14 Center exposures, and I want to just make a
15 disclaimer that this is totally outside of my area
16 of expertise so the metals exposure levels are very
17 complex in World Trade Center. And I tried to, in
18 the next few slides, give you a sense for some of
19 the concerns but I don't have any kind of a
20 conclusion to the extent that I did for VOC.

21 So on the next slide, Cahill and colleagues have
22 thought a great deal about the metals and other
23 exposures generated at the World Trade Center site,
24 and they've developed an incinerator hypothesis
25 which provides an explanation for the very fine

1 aerosols that were liberated. And a number -- and
2 just basically it would be the temperature that
3 would be involved in these very fine aerosols and
4 there were, his quote, unprecedented levels of
5 several metals. Also, his quote, and this again is
6 from the very fine aerosol chapter in the American
7 Cancer Society book that Liz had referred us to,
8 he's commented that the health concerns focus on
9 workers at the site, as plume lofting protected
10 most of New York City. What I don't know in that
11 regard is the impact on residential -- residences
12 that were very near the site. I know others have
13 commented this afternoon on high rises that were
14 right near the site, so that's something to think
15 about.

16 And the next slide, he comments that some metals,
17 and lists a series occurring at unprecedented
18 levels in these very fine aerosols, and then goes
19 on to note that levels dropped off dramatically,
20 even over the course of the month of October and
21 definitely by the end of May.

22 There are other slides listing a variety of metals
23 that have been found both in dust, but the concern
24 that dust is present after the fact may not be
25 representative of what people actually breathed in

1 at the time. I'm told indicating that lead levels
2 do not appear to be a huge concern.

3 Skipping to the next slide, Liroy's comment. The
4 concern that deposited material with metals in it
5 could lead to ongoing exposure -- because in
6 contrast to VOCs, metals are very persistent in the
7 environment. Liroy commented that concentrations of
8 arsenic and cadmium were relatively low but still
9 in the parts per million range, so we need to keep
10 that in mind when thinking about dust.

11 Next slide, a little bit of data, some of the small
12 amounts that I found regarding airborne levels
13 other than in the plume.

14 And then finally metal implications. So the metals
15 data are hard for me to synthesize in terms of
16 thinking about risk to individual workers. There's
17 been a lot of characterization of the plume, and
18 I'm not up to speed on all of it at this point, but
19 the thoughts that I have in terms of the metals at
20 this point are the potential risk for toddlers who
21 spend a lot of time on the floor and do a lot of
22 hand to mouth activities from persistent metals in
23 dust in residential areas. And then my other
24 concern is the impact that these metals in dust,
25 these very small particles, being deposited in the

1 lungs, and I'm wondering, you know, some of these
2 metals do bioaccumulate. We, you know, lead and
3 cadmium clearly reside in the body and accumulate
4 but I'm wondering if that very high initial load
5 could change the half-life of some of these metals
6 in the body, and I'm also wondering about the
7 potential for interaction with the very high pH,
8 although I don't know that if some materials that I
9 read commenting that the smaller particle size had
10 a more neutral pH, so I don't know how significant
11 that concern is. But I did want to mention that.
12 So that's all I have.

13 DR. WARD: Thank you. Where do we stand on time,
14 Paul?

15 DR. MIDDENDORF: We need to get started.

16 DR. WARD: Okay. We're going to start public
17 comments now and then we'll get back to Virginia
18 with any questions.

19 **PUBLIC COMMENTS**

20 DR. MIDDENDORF: Okay, each of our public
21 commenters has signed up on a first-come-first-
22 serve basis, and each of them will have up to five
23 minutes to present. I remind people that it's
24 often surprising how quickly five minutes can go
25 when they talk about a subject of great importance

1 to you so when you reach four minutes, I'll let the
2 commenter know that they have one minute remaining,
3 so they can be sure to make the points that they
4 want to make in that last minute they have.

5 If they get up to five minutes, I'll have to rudely
6 interrupt them and thank them for their comments.

7 I apologize up front to anyone to whom that happens
8 but we have to be fair to all of our commenters.

9 We do have one commenter this afternoon who will be
10 on the phone, and just remind them to keep the
11 phone on mute until I call out their name, and then
12 they can unmute the phone and they'll have the same
13 five minutes everyone else does.

14 Also want to point out that everyone has the option
15 of submitting written comments to the docket for
16 this committee. The docket number is 248, and
17 information on how to submit comments is in the
18 Federal Register Notice; it's also in the NIOSH
19 docket page, and it should be on our committee web
20 page as well.

21 Lastly, I want to remind our commenters of the
22 redaction policy for public comments. The policy
23 is stated in the Federal Register Notice for this
24 meeting; it's also on the committee's web page and
25 it's posted at the registration table if anybody

1 wants to look at it. And the policy outlines what
2 information will be kept and what information will
3 be redacted before it's posted to the docket.

4 So when I call your name if you would kindly come
5 up to the podium. We need to get the microphone up
6 there, wherever it is, handheld mic? Our first
7 speaker is Micki Siegel de Hernandez.

8 MICKI SIEGEL DE HERNANDEZ: Good afternoon. My
9 name is Micki Siegel de Hernandez. I'm the Health
10 and Safety Director for the Communications Workers
11 of America in District 1. Our union represents
12 several different groups of 9/11 responders as well
13 as area workers affected by 9/11 exposures. I'm
14 one of the designated labor reps on the World Trade
15 Center Health Program Responder Steering Committee
16 and a member of the World Trade Center Health
17 Program Survivor Steering Committee and was the
18 sole labor liaison for the EPA World Trade Center
19 Expert Technical Review Panel.

20 First, regarding adding cancer to the list of World
21 Trade Center-covered conditions, our union supports
22 that. The time is now and I believe that today's
23 presentations, thankfully, provide ample support
24 and rationale.

25 Secondly, regarding the research agenda topics, it

1 was good to see such a breadth of topics suggested
2 by the STAC. We support research on cancer, heart
3 disease and other chronic conditions, mechanisms of
4 inflammation and disease persistence which could
5 hopefully lead to more effective treatments,
6 immunological disorders including autoimmune
7 conditions and nervous system disorders.

8 We would also like community-based participatory
9 research projects involving affected responders,
10 area workers and residents to be encouraged.

11 While funded research is important, it can't be the
12 sole source of our understanding of World Trade
13 Center-related disease, and I cannot emphasize
14 enough the need for improved and continuous disease
15 surveill -- disease and symptom surveillance in the
16 World Trade Center Health Program. This deserves a
17 closer look.

18 A couple of examples are headaches, loss of
19 peripheral vision, symptoms which are nonspecific
20 and can have many causes but are frequently
21 described by responders. While aerodigestive
22 disorders may be the most common World Trade
23 Center-related conditions, they are not the only
24 ones. However, if you are not looking for other
25 illnesses, you will never find them.

1 And then I have some sort of random comments that
2 were taken from the presentations today regarding
3 exposures. First, in several presentations it was
4 mentioned that there were no samples that were
5 taken during that critical first week after the
6 World Trade Center collapse. I think that needs to
7 be revised to say that no measurements were
8 reported rather than none taken.

9 In a joint statement of the EPA and OSHA on 9/14,
10 they stated that sampling data for asbestos were
11 below levels of concern, not likely to cause
12 long-term health effects. Christie Whitman's
13 famous statement on 9/17, declared the air and
14 water safe based on initial sampling. EPA pulled
15 early sampling data from their website, the New
16 York City Department in Environmental Protection
17 hazmat team was onsite that first day, took samples
18 that were never reported.

19 So this is indicative of a stance taken by
20 government agencies that they have stuck to to this
21 day, and in part explains the disconnect between
22 reported sampling, or non-reports, and actual
23 health effects.

24 It also, as was discussed in several of the
25 presentations today, it matters what you sample

1 for, when you sample, where you sample, how you
2 sample and how samples are analyzed.

3 This also explains in part the inconsistency with
4 levels being reported as safe and the health
5 effects. Sampling was not conducted in a
6 consistent or even comparable way. It was done by
7 several different agencies, much of the sampling
8 was done by private entities and therefore not in
9 the public record.

10 I would also argue that a wrong model was used.
11 Individual contaminants were measured when the
12 World Trade Center dust and fire, the plume from
13 the fire, is a very complex mixture. There were
14 different standards that were applied that were not
15 health-based standards, and these were used to make
16 statements about health; such as the OSHA
17 standards. The PELs are not health standards and
18 they are also based on 1960s science and knowledge.
19 Ambient air exposures are also but one part of an
20 individual's exposure. In some of the articles,
21 there was an article that was distributed about,
22 the Liroy article, about environmental conditions
23 and human exposures at a current
24 post-September 11th, 2001, in 2006, --

25 DR. MIDDENDORF: You have one minute left.

1 MICKI SIEGEL DE HERNANDEZ: One minute? And in
2 that it said that the second rain event washed much
3 but not all of the remaining outside settled dust
4 and smoke away; this is simply not true.

5 Lastly, the duration of exposures were short-term
6 for many people. This was repeated in a couple of
7 presentations, the committee should be careful
8 about how it defines or thinks about short-term
9 exposure, what is known and not known about
10 exposures.

11 Is it short-term for responders working up to eight
12 months at Ground Zero for 10- to 16-hour or more
13 shifts? Is it short-term for responders who
14 continued response and restoration activities in
15 contaminated areas well after the site was closed?
16 And you should also know that there is no known end
17 date for any given individual or for areas since
18 levels of contamination and exposures, particularly
19 in indoor sites, were not assessed. Thank you.

20 DR. MIDDENDORF: Our next speaker is Bruce Edwards.

21 BRUCE EDWARDS: Thank you for giving us the
22 opportunity to speak at this meeting. My name is
23 Bruce Edwards. I am a permanently disabled IBEW
24 Local 3 journeyman electrician. I was asked to
25 work at the Verizon building at 140 West Street.

1 The building is across Vesey Street from where the
2 North Tower and Building 5 stood. 140 West Street
3 was severely damaged by falling debris of the
4 towers on its south side and the collapse of
5 Building 7 to its east.

6 I arrived at Ground Zero early in the morning of
7 September 14th. Our arrival at the site was delayed
8 due to fear of instability at the site, and we were
9 originally scheduled to arrive the previous day.

10 I was employed by an electrical contractor that was
11 known as a Telco contractor, very knowledgeable in
12 the operations of telephone central offices. We
13 were tasked with the temporary restoration of
14 electrical power by means of portable generators.
15 The reason this work was so important was due to
16 the antiquated underground cabling methods of
17 downtown Manhattan. The Verizon building at 140
18 West Street was the main path of communications in
19 and out of the Wall Street business district, and
20 most importantly, the New York Stock Exchange.

21 The president at the time, George Bush, had ordered
22 Verizon to restore communications as soon as
23 possible. Due to our efforts, the Stock Exchange
24 was up and running on Monday September 17th, before
25 the opening bell.

1 We continued working at 140 West to permanize (sic)
2 the temporary work to safety and then actually
3 repair the building. It was many weeks before Con
4 Ed could get power to the area at Seven World Trade
5 Center, was the substation, the power substation,
6 of the area. Our portable generators were needed
7 to operate the building.

8 In the first few weeks, we worked 16 to 18 hours
9 per day, seven days a week. And then as our
10 numbers increased, we went to two shifts, 24 hours
11 a day. As a supervisor, my responsibility extended
12 to both shifts.

13 I'm sorry about all the background but I believe
14 that is important to understand that the reason
15 that I was asked to work there, and believe me, you
16 didn't have to ask me twice. I felt a bond to the
17 World Trade Center, as my father and brother had
18 both worked on the construction, and we had been
19 attacked. Nationalism and patriotism was at an all
20 time high.

21 Ultimately though, I was a civilian required --
22 requested to work in a disaster area with little
23 protection and no knowledge of the long-term
24 problems that could occur. My original crew on the
25 first day consisted of myself and seven other

1 electricians, basically an advanced team to lay the
2 groundwork. Within a few days, we had well over a
3 hundred electricians on site.

4 Now, if you ask me would I do it again, my first
5 instinct is yes. Like many, I took this
6 personally. But in further review, I'm afraid I
7 might not do this because the price I paid was
8 steep. In April 2007, I was diagnosed with
9 stage IV, non-Hodgkin's lymphoma.

10 I spent nearly two years in and out of hospitals
11 for chemotherapy treatments, and fortunately I was
12 able to have a stem cell transplant in
13 December 2008. I'm currently in remission but
14 remission isn't a cure. I live with the constant
15 thought that the next low-grade fever I get is a
16 return of my disease.

17 But even then I consider myself lucky because of
18 the original eight, Robert Kiano (ph) didn't fair
19 as well. He succumbed to his disease in 2010 at
20 the age of 50. I was 50 when I was diagnosed also.
21 Now I'm no scientist but I do see of our original
22 crew two cancers out of eight. That's a 25-percent
23 disease rate in relatively young men.

24 I was forced to retire from my career at least ten
25 years early. The financial hit was crippling. I

1 had two children in college and practically no
2 money flowing in.

3 The next problem was clinical depression from all
4 the problems there. Fortunately, with some good
5 doctors, I was able to clear that.

6 DR. MIDDENDORF: One minute, please.

7 BRUCE EDWARDS: In the time since 9/11, some
8 troubling items have emerged. Our government seems
9 to have downplayed, and I use the term graciously,
10 some of the conditions at Ground Zero.

11 Ms. Whitman's the air is safe declaration and the
12 release of some information about the accident
13 exposure. The report released around the tenth
14 anniversary showed dioxin levels 1,000 times higher
15 than normal, and the highest the EPA has seen.
16 What is especially troubling is the sampling began
17 on September 23rd. That's almost two weeks after
18 the attack.

19 The next two months the sampling continued and
20 showed steady decline, so I can only imagine what
21 the levels were on day one, or day four for my
22 crew.

23 The report from the fire department is also an
24 eye-opener. Here's a segment of the population
25 that is generally in good physical condition and

1 well-monitored, and yet the cancer levels for those
2 exposed at Ground Zero is well above normal.

3 What I have come to learn is that --

4 DR. MIDDENDORF: Your time is up --

5 BRUCE EDWARDS: Okay. Well.

6 UNIDENTIFIED SPEAKER: Let him speak.

7 BRUCE EDWARDS: I'd just like to let people know
8 here that the cancer rates are very high for a
9 young population where normally they would be in an
10 older group. And I implore you to add cancer to
11 the bill as the Senate, I should say the Congress,
12 has done with this letter that they sent to you.
13 Thank you.

14 DR. MIDDENDORF: Our next commenter is on the
15 phone. Rich Dambakly. If you would unmute and
16 begin your presentation.

17 RICH DAMBAKLY: Hello?

18 DR. MIDDENDORF: We can hear you.

19 RICH DAMBAKLY: Okay. My name is Richard Dambakly.
20 I'm an underground worker for Verizon, at least I
21 was an underground worker for Verizon. I worked at
22 Ground Zero from the moment of the disaster, every
23 day for six months straight, 12 to 16 hours a day,
24 no days off.

25 I developed the World Trade Center cough. And for

1 those of you that are unaware what this feels like,
2 it's a cough where your chest is exploding out of
3 your body that doesn't stop.

4 In March of 2002, it had gotten so bad I had to go
5 to emergency. After being diagnosed with lymphoma
6 cancer, I started intense chemotherapy treatment
7 that lasted five months.

8 Just recently someone mentioned to me that the
9 actor Andy Whitfield from the television show
10 Spartacus had died from lymphoma, and it was his
11 second occurrence. And here I am with no CAT scan
12 for three years because I have -- I can't afford
13 one. I have no medical insurance. How do you
14 think that makes me feel?

15 I'm a father of five children, my oldest being 15.
16 My family needs me. I want to be around to walk my
17 daughters down the aisle and play ball with my son.
18 Should I become a beggar and maybe raise the money
19 for a CAT scan? Just like our Vietnam vets, that
20 they were forgotten?

21 So many have died already from cancer. Their
22 families need help now. This can't go on. When
23 other countries are in need, we don't waste a
24 minute. Immediately we send them money. We ask
25 for nothing in return. When President Bush arrived

1 at Ground Zero, I stood and listened to him speak
2 to us and tell us to stay strong, stay here, help
3 us, do whatever it takes, whatever you have to do,
4 work any amount of hours. We need you; we'll be
5 there for you. And we did it, each and every one
6 of us that stayed strong. Anything we could do in
7 our power. No one said, I can't help or that's not
8 in my job description. No, we did whatever we were
9 asked and more. The country needed us and that's
10 all that mattered.

11 So now that we need the help and when you should be
12 strong for us, instead you're taking the position
13 that covering us for cancer is not in your job
14 description, and that's wrong.

15 On 9/11 terrorists came to our country and were
16 responsible for thousands of deaths. Don't give
17 them more reason to celebrate by not responding to
18 our country's aid and causing more American lives.
19 Don't allow them more victory than they already
20 have.

21 We were there when our country needed us, and our
22 country should be there for us when we need them.
23 God bless all my fellows and other survivors and
24 first workers in the World Trade. God bless you
25 all. Thank you very much.

1 DR. MIDDENDORF: Thank you, Mr. Dambakly.

2 Our next commenter is Alex Sanchez.

3 ALEX SANCHEZ: Good afternoon to members of the
4 committee; my name is Alex Sanchez. This good? I
5 am a 9/11 responder, clean-up worker. On
6 September 11th I had a very close encounter with
7 terror. I was standing not very far from where
8 this building is today.

9 On September 13th to March 15, I performed cleanup
10 with other cleanup workers in the skyscrapers
11 surrounding the pit. Ten buildings in a period of
12 six months. Twelve-hour days, seven days a week.
13 Some of the buildings I worked in included 1, 2, 3
14 World Financial Center. I had a ringside seat to
15 what police officers, firefighters were doing at
16 Ground Zero. When I went past those barricades, as
17 a citizen, as a New Yorker, I knew what was
18 expected of me.

19 When men and women started getting sick and dying,
20 I also knew what was expected of me. Since late
21 2003, early 2004, I've been walking the halls of
22 Congress alongside many of the men and women who
23 are in this committee and who are also here today.
24 John Feal, my mentor, president of the FealGood
25 Foundation, an officer and a gentleman,

1 paratrooper, United States Army. We do not leave
2 ours behind. What message are we sending to future
3 generations and to the international community when
4 we overlook and not appreciate the work and the
5 efforts of those who served at Ground Zero?

6 Let me give you some facts. Basically you should
7 know these by now. Seventy percent of the men and
8 women who came to Ground Zero are suffering from
9 lung disease, chronic gastric disease, post
10 traumatic stress disorder. I'll give you another
11 example.

12 Jim Ryan, John McNamara. Both on the same office,
13 Senator Lieberman, two months later, I asked my
14 assistant director, Monroe Checko (ph), who is this
15 gentleman? John McNamara disintegrated in a period
16 of two months.

17 We don't need bigger government or smaller
18 government. What we need is responsible
19 government, government that takes care of the
20 people. Enforce and enact laws, current laws. I
21 am a single father of an amazing 10-year-old. This
22 is not the message I want to send to my son, my
23 country cannot get it right. Ten years down the
24 road cancers are killing the men and women who came
25 to Ground Zero. Exposure science tells us that

1 when you are exposed to high level of toxicity, you
2 need 15 to 25 years of medical treatment. We only
3 got five. We cannot continue to play games with
4 human lives. We need to stand up. We need to
5 serve those who serve our country. We shall never
6 forget and may God bless the United States of
7 America. Thank you.

8 DR. MIDDENDORF: Thank you, Mr. Sanchez. Our next
9 commenter is John Feal.

10 JOHN FEAL: How's everybody doing today? Good? I
11 don't think I need a microphone. I'll introduce
12 myself when I'm done. This way I can get my five
13 minutes in.

14 One, I want to thank NIOSH for doing this. I want
15 to thank the STAC committee for hearing me today.
16 I'm not here to ask you to add cancer to the bill.
17 I'm here to ask you add certain cancers to the
18 bill. I'm getting a little tired of hearing we
19 need to add cancer to the bill. You cannot add
20 every cancer to this bill; that's impossible. I
21 get it. I worked on this bill for eight years,
22 more than most people in this room. But there are
23 cancers, unequivocally, undoubtedly, that need to
24 be added to this bill yesterday.

25 I am never the smartest man in the room and I'm not

1 even the smartest man at this podium probably, but
2 it doesn't take a scientist or a doctor to know
3 that 9/11 and its toxins have caused these blood
4 cancers.

5 For years when we walked the halls of Congress, we
6 were applauded for the way we approached Congress
7 to get this bill passed. And when we were lobbying
8 to get that bill passed, we were lobbying to get
9 cancer added to that bill. But during the
10 negotiations, that was taken from us. But I am
11 going to use the same zest and the same energy to
12 help get those certain cancers added to this bill.

13 I will occupy Ground Zero. Don't worry about
14 Occupy Wall Street. I will do whatever it takes
15 because at the end of the day, I care about human
16 life. I don't care about what you're having for
17 dinner, I don't want to go to your house for
18 coffee. I care about human life. I care about
19 adding cancer, certain cancers, to this bill.

20 And as for epidemiology, let that not be your only
21 role model. Epidemiology can only do so much, like
22 the cancers that we know that should be added, use
23 epidemiology on that. 9/11's unprecedented. It
24 never happened before. So use something else other
25 than an epidemiology. And believe me, I can't even

1 spell the word, that's how smart I am not. Okay?
2 So I'm asking you guys, with power comes
3 responsibility. You have a responsibility today,
4 tomorrow and from this day forward to do what is
5 morally right.

6 I just came from a press conference at City Hall,
7 and I almost threw up on myself listening to people
8 who do not know what they're talking about. But
9 appreciate the magnitude of this 'cause I do. I
10 lost half a foot ten years ago. Eleven weeks in
11 the hospital. I'm lucky but I feel guilty that I
12 can go to Sheelar (ph) and say I want to apply for
13 the Zadroga bill 'cause I lost half my foot.

14 Boohoo. Say that to John Walcott or Arthur Noonan,
15 who have leukemia and blood cancers. That should
16 be added yesterday. You're playing God right now.
17 Our fate is in your hands.

18 I am the nicest guy in the world. I want to be
19 your friends. But like I told every member of
20 Congress and every member of the Senate when I met
21 them for eight years with this bill, I will do
22 whatever it takes to get cancer added to this bill.
23 Thank you.

24 DR. MIDDENDORF: The document which you handed out
25 to the committee members will be added part of the

1 docket. Just wanted to let you know that but it
2 may be redacted to some extent. We'll have to look
3 further.

4 JOHN FEAL: Do what you please with it.

5 DR. MIDDENDORF: Okay, our next commenter is T.J.
6 Gilmartin.

7 T.J. GILMARTIN: Good afternoon. My name is T.J.
8 Gilmartin, and I'm 32 years as a foreman and a shop
9 steward building high rises in New York City with
10 the union.

11 Now, I had to go to so many OSHA classes for these
12 high rises of stuff they taught us was cancerous
13 and, you know, don't do this, don't do that.
14 Everything, everything I been taught to and told is
15 dangerous and cancer-causing is being thrown out
16 the window on this World Trade Center. I mean, I
17 know what goes into building a high rise and one
18 thing that was -- and the Trade Center was built
19 prior to 1973, when the asbestos was in the pipes,
20 it was in the cement, it was the silicosis, the
21 heavy metals, the chemicals and the PCBs.

22 Does anybody know about those electrical vaults in
23 the basements of those trade centers? You know
24 that's totally cancer-causing chemicals inside
25 those -- the vaults and the transformers? Okay?

1 All that was there and we never hear of anything.
2 Anything about any of that.

3 I mean, all this stuff is concern -- is confirmed
4 as a federal cancer-causing chemicals. The
5 building was totally filled with all these
6 chemicals. The fire department, the PDA have done
7 studies showing that their men are dying a lot more
8 than they are usually dying fighting fires.

9 I mean, OSHA would lock me up if I was -- if I was
10 grinding concrete on a high rise and that powder,
11 if I didn't have a battery-operated respirator, I'd
12 be locked up by OSHA, either thrown in jail or
13 fined for having my men do that. I mean, you had
14 220 stories of pulverized concrete besides
15 everything else that, God forbid, was going to
16 happen in another nine years with the asbestos,
17 with that 20-year lag time.

18 It's been over ten years since the World Trade
19 Center was destroyed, and that's been a time so
20 many first responders have paid with their lives.
21 The percentage is out of whack compared to how many
22 first responders just tried to help their fellow
23 man. It seems to me that this is all about the
24 money. I mean, I understand that you'll have
25 everybody claiming that they got cancer from World

1 Trade Center but like John said, there were certain
2 cancers from the ears, nose, -- I mean, your mouth,
3 your nose or absorption that should be covered by
4 this.

5 But it's -- you know, I mean, that's basically what
6 I have to say. I mean, just that I been in the
7 business of high rises and I know what causes
8 cancer on these things and, you know, you put up a
9 high rise, OSHA's there, you're doing it, you know,
10 you're in a lot of trouble if you do it that way.
11 Everything that could get you cancer on a new high
12 rise was all down at the Trade Center, and it was a
13 lot worse because it was built before 1973 when the
14 world was changed. Thank you.

15 DR. MIDDENDORF: Thank you very much,
16 Mr. Gilmartin.

17 Our next commenter is Thomas Fay.

18 THOMAS FAY: Good afternoon, ladies and gentlemen.
19 Is this the speaker here? My name is Thomas Fay,
20 and I come from a town at the Jersey shore called
21 Spring Lake, New Jersey. On September 11th I was
22 getting my wisdom teeth pulled; and the planes hit
23 the building and I raced home and proceeded to
24 watch on television for about 36 hours. And after
25 the 36 hours, I couldn't take it anymore so being a

1 volunteer fireman for over 37 years in the Spring
2 Lake fire company in Spring Lake, New Jersey, I
3 decided to go get my gear, jump in my car and race
4 to New York. I got there in 50 minutes, which is
5 unprecedented.

6 I was directed down to the south end of the city
7 and parked my car on 14th Street and I walked in.
8 Two other firemen drove by this desolated area of
9 lower Manhattan and picked me up. I never knew
10 them before but I know them now. Both are very
11 sick.

12 They drove me down and they went out to get a
13 camera that day to take pictures. I didn't want
14 any pictures taken of me that day; I was there to
15 work, not to have any pictures taken. But lo and
16 behold, they took two pictures of me and those two
17 pictures ended up being the proof that I needed to
18 show that I was there.

19 The disease that I contracted from my 12 hours
20 working on the south tower pile, solely on
21 September 13th, was non-Hodgkin's lymphoma,
22 stage II, B-cell aggressive. The way that was
23 found in me was that I, in 2007, after the
24 disaster, a friend advised me that I needed to go
25 get checked out at the World Trade Center medical

1 monitoring treatment program they had at Rutgers,
2 which I did.

3 I went in 2007, 2008, and in 2009, I noticed a lump
4 in my left leg. I showed it to Dr. Iris Udasin out
5 there. She said you've got to go to New York City,
6 Mt. Sinai immediately. Within a week the tumor was
7 taken out. Four days later I was told that I have
8 cancer.

9 I fought the battle brave and hard. I'm in
10 remission now which is a good thing, but for people
11 like us that went up there and put our time in, I
12 being a volunteer, I was paid nothing, I would go
13 again tomorrow because of one thing: I love my
14 country. That's it, pure and simple.

15 Being a guy from the Jersey shore, a popular person
16 everyone knows who comes from down there is Bruce
17 Springsteen. He has a new album out. And he has a
18 song on it called, *We Take Care of Our Own*. That's
19 the theme song for us first responders. We want
20 our government to take care of us.

21 We went in there. We fought hard. I worked 12
22 hours on that burning pile. If I fell once, I
23 would have been cut to shreds. But that wasn't on
24 my mind that day. On my mind that day was to help
25 as many people as I could. That's why I joined the

1 fire department, to help people. I didn't join the
2 fire department to get cancer.

3 My cancer's in remission but as of Monday, a recent
4 trip to the doctor, has shown that I now have skin
5 cancer. I'll fight that battle on my own and take
6 care of that as I should. But it is my hope that
7 this -- people here, grouped here today, do the
8 right thing, which is to include blood cancers in
9 the Zadroga bill. Thank you very much for your
10 time.

11 DR. MIDDENDORF: Thank you very much, Mr. Fay. Our
12 next commenter is Arthur Noonan.

13 ARTHUR NOONAN: Hello. My name is Arthur Noonan,
14 retired now but back in September 17th, 2011, I was
15 employed by the Chicago Fire Department. As the
16 last speaker, we were watching on television
17 nonstop at the firehouse. Finally we couldn't take
18 it anymore, we saw what a devastating effect this
19 had on the country as well as to New York, and we
20 decided to come here. I believe there was a group
21 of 14 of us. We flew in and we spent seven days
22 working here.

23 I was a pretty healthy guy as well as the rest of
24 the people that came with me. A lot of young
25 firemen from Chicago, good firemen, and we did

1 everything from cleaning tools and changing blades
2 and batteries in the tool shed, until we finally
3 got to work on the actual Pile. Some days we would
4 cut aluminum off of steel beams so the iron workers
5 could cut the beams in sizes small enough to fit on
6 the trucks to haul them away.

7 Eventually we got to work on the Pile. You'd start
8 at the back of the Pile, there might be a hundred
9 firemen in front of you. You'd pass buckets
10 forward empty, and backwards full. Finally you'd
11 get up to the point where you were the one that was
12 digging. You'd be on your hands and knees; what
13 respirators we had didn't work, they kept clogging
14 up or from the sweat would just turn like a mud on
15 there. We finally had to take those off. But you
16 kept working because you knew your brother
17 firefighters, policemen and many loved ones of
18 civilians who were also in that Pile. And all we
19 wanted to do was try to close a part of life for a
20 lot of people.

21 In December 2004, I became ill at work, was taken
22 to the hospital. Thought I had a bad touch of the
23 flu; everyone was sick in the firehouse then. It
24 was the day before Christmas Eve. They let me go
25 home for Christmas Eve and Christmas Day, I had to

1 come back the following week, and I was diagnosed
2 with AML, acute myelogenous leukemia.

3 I went from 210 pounds to about 140 pounds in six
4 months, had several chemo treatments, and luckily I
5 am now in remission. But remission is not getting
6 better. It just means they're holding you steady
7 so every day you hear something on the radio,
8 whether it be a celebrity or sports figure, just
9 recently we had a famous singer die of leukemia.
10 Every time you hear that word leukemia, it all
11 comes back to you.

12 When we came to New York, we did it on our own. We
13 did not expect to get anything for it. We just
14 wanted to help our country. We wanted to show the
15 world the support that New York and the United
16 States, how they all come together in a time of
17 need.

18 Personally I have taken a tremendous loss on my
19 medical benefits. I've gone through about three-
20 quarters of what I'm entitled to in my lifetime for
21 myself and my wife and if this comes back, I
22 probably only have a few hundred thousand dollars
23 left in my medical plan from the City for
24 treatment. After that, I don't know what I'll do.
25 So I'm hoping that cancers, certain cancers, will

1 be included in this so people that came to help do
2 not have to have that constant worry in their mind
3 if their cancer comes back, they won't be able to
4 get any treatment. Thank you.

5 DR. MIDDENDORF: Thank you very much. John
6 Walcott.

7 JOHN WALCOTT: Hi. My name is retired detective
8 John Walcott. Like everyone else here, I'd like to
9 thank you for this opportunity.

10 I also was diagnosed at 38 with AML leukemia. As I
11 stand here in front of you I've had six months of
12 chemotherapy, stem cell transplant, and I have
13 other illnesses that are recognized in the Zadroga
14 Act. But looks are deceiving. All my nerve
15 endings are burnt out all my -- in my hands and my
16 feet. There's not a day that goes by I'm not in
17 constant pain.

18 The City retired me due to my leukemia, which they
19 said I got from 9/11. Social Security recognized
20 it. It seems that only the country doesn't
21 recognize it.

22 Before 9/11, I was approximately 36 years old. I
23 was never sick a day in my life except for the
24 common cold. I was a very extremely active
25 narcotics detective, well over 3500 arrests in my

1 career involved in. I was a high school hockey
2 coach. Used to do physical activity, lift, run
3 every day. No longer can do any of that. I was on
4 the fast track to probably becoming a hockey coach
5 in college. We had an exceptional team,
6 exceptional record and I turned down many jobs
7 which I planned to take when I retired. Which,
8 that's been cut short.

9 On 9/11 itself I wasn't scheduled to work 'til late
10 that evening. I was told what happened, I was
11 woken up, and I was down there in 93. So without
12 hesitation, I ran right down there to help my
13 fellow detectives or policemen at the time.
14 Shortly after the second tower had collapsed, I
15 arrived.

16 Did -- from recovering bodies, body parts, to Mayor
17 Giuliani even assigned us one day to VIP tours for
18 all his friends. So I've done everything, cut
19 steel. You weren't a policeman when you were down
20 there; you were just somebody trying to help.
21 As I told you before I had the transplant and
22 everything else.

23 Well, you know, let's talk a little bit why we're
24 down here. We all know that the benzene and
25 asbestos and all over cancer carcinogens were down

1 there. That's no secret. I mean, that's been for
2 a hundred years. We don't know what they do if you
3 mix them all together nor do I think anybody really
4 cares because if they did, it wouldn't have taken
5 us ten years to get to this point.

6 We know there's a usually high number of early
7 responders that are diagnosed with cancer. Yet no
8 one seems particularly interested in trying to
9 corroborate any of these findings at the site, at
10 the cancer rate. The large population of
11 responders and workers are being looked at, which I
12 think you guys are doing a study of over 50,000
13 people. But I think that study's wrong. I think
14 you should study guys and girls and everybody who
15 was down there the first day, first week, first
16 month. And if we do that, you're going to see that
17 the 362 PBA Study, that rate is going to be
18 astronomical. It's probably going to be in your
19 60s to 70 percent of cancer rate.

20 There's many reasons. We all know there's many
21 reasons why the City's and the country's not
22 releasing these numbers. Because they're doing you
23 a 50,000 population rather than a 2500 to 5,000
24 population. So that statistics are going to be
25 extremely less and it's not going to prove cancer.

1 But if you did, if there was actually 2500 to 4,000
2 that were down there the first week, day or month,
3 it's going to be astronomical. And then the red
4 flag is going to be up.

5 But when there's litigation going on and there's
6 hearings about to happen, what do we do? We have
7 to make the numbers look bad because the City kind
8 of painted themselves in a corner right now with
9 this.

10 DR. MIDDENDORF: One minute left, please.

11 JOHN WALCOTT: Okay. You know, I think that's
12 where we need to concentrate. We have to
13 concentrate on -- let's concentrate on 2500 to the
14 3,000 that were down there versus that. I don't --
15 there's a part of me that envies you folks and
16 there's a part of me that doesn't envy you folks.
17 You have to make a tough decision. But luckily for
18 you folks you have ten years and weeks of hearings
19 to make this decision.

20 I had a phone call and I had to rush down. Now I'm
21 sick, my daughter'll never see me walk her down the
22 aisle. I can put my head on my pillow and go to
23 sleep at night knowing I did something that in the
24 recovery that meant closure for people. You folks
25 have that same power now. Twenty years from now if

1 the cancer isn't added, and my grandchildren, that
2 I'll never see or hear, do you say you made the
3 right mistake? Did you make the right decision?
4 Thank you.

5 DR. MIDDENDORF: The next commenter is Reginald
6 Hilaire.

7 REGINALD HILAIRE: Hi. Good afternoon. I'm a
8 police officer with the NYPD for 11 years. I was a
9 rookie when 9/11 happened. I'm currently assigned
10 to PSA 5, which is a housing precinct up in East
11 Harlem. I worked over 850 hours combined at the
12 World Trade Center and Sandman Landfill.

13 In 2005, shortly after my son was born I was
14 diagnosed with thyroid cancer. I immediately asked
15 my primary care physician if this was related. He
16 said, he looks at my lump and said, what were you
17 exposed to down there? I've seen him since 1999,
18 before I became a cop. So 2005, I had total
19 thyroidectomy, radiation and ever since then I take
20 a pill, a synthroid, and it regulates my thyroid.
21 Winter of 2005, I go back to my primary care
22 physician, he noticed my blood count was pretty
23 low. He refers me to a hematologist and that
24 hematologist does a bone marrow biopsy, and he
25 comes back and he says, the pathology report -- I

1 disagree with the pathology because it says you
2 have multiple myeloma but I disagree. You're too
3 young to have this. He repeats it in 2006, it
4 comes back multiple myeloma. He's still confused.
5 I go -- I sent everything to Sloan-Kettering. They
6 do another biopsy, bone marrow biopsy, April 2006.
7 They confirmed it. I thought okay, great, treat
8 it. No, we can't treat you because you have
9 smoldering multiple myeloma, early stages. So I'm
10 like, is there anything out there for me? No, you
11 can't -- there's nothing. We have to wait until it
12 gets worse in order to treat you. He says within
13 two to three years, you have 50, 60-percent chance
14 of it getting worse.
15 Thankfully every four months now I go to
16 Sloan-Kettering, they do blood work, urine work,
17 and if I get the phone call, that means it's not
18 good. So far, knock on wood, everything's okay.
19 I have no family history of cancer. I'm pretty
20 much the healthiest one. I am a son of Haitian
21 immigrants. I am the only member of my family
22 that's a police officer. I was born and raised
23 here, still work here in Harlem. I can't retire
24 because, even though I'm not really sure if I want
25 to, but I can't retire because I'm not sick enough

1 so it's an oxymoron right there.

2 I have two red cancers. I don't -- I work with a
3 lot of cops in PSA 5. I don't know why I have it.
4 It's just one of those things I've come to accept
5 it. In 2006 I read an article in the Post saying
6 that there's other first responders with cancer. I
7 contacted that reporter who introduced me to one
8 detective who has lymphoma. He introduces me to
9 others. I got to know about 11, and I'm pretty
10 close to about four of them. Three of them have
11 multiple myeloma. I never met them before in my
12 life.

13 I met one police officer through the PDA who
14 (unintelligible) I did. His name was Bob Rossalein
15 (ph); he had (unintelligible) cancer. We got to
16 talk for about a year and then he eventually died
17 in 2010. So I always think about him, think about
18 his family, I'm still close to his widow.

19 I don't -- I'm not a scientist; I'm just a cop, I
20 just want to do my job. I think a lot of us want
21 to do our jobs. I don't think it's coincidence. I
22 never met these people before in my life.

23 Someone asked me before if they had to do it again.
24 I, like I said, I'm still with the NYPD. I'm doing
25 clerical work. I'm pretty now senior now. If it

1 happens, again, and I'm pretty sure it would, would
2 I do it again? Would I tell my junior cops to go?
3 I don't know. I love New York City, I love the
4 people here. I'm not fond of the government. They
5 showed so careless without a doubt.

6 What's really insulting, I could deal with cancer,
7 I could deal with questions, how you doing. As a
8 New Yorker, how you doing could mean ten different
9 things. How you doing or in my case, so how are
10 you doing?

11 DR. MIDDENDORF: One minute, please.

12 REGINALD HILAIRE: What I can't stand is
13 politicians, everybody can say, okay, great, great
14 job; you're heroes but when it comes to treating
15 us, hold back. It's just too early to step up the
16 study; it's not there yet.

17 I try to tell the cops in my precinct get yourself
18 checked out. They look at me. We can handle
19 perps, we can handle perps with guns, we can even
20 handle bosses that are rough. We can't handle our
21 own mortality.

22 So I urge all of you, just like us, when they call
23 us heroes, all of you can be heroes by just saying,
24 adding cancer. You will save lives by putting
25 cancer in the bill because it will tell first

1 responders to get checked out. You don't know how
2 much of a difference you guys will make if you add
3 cancers. You will tell somebody with the public --
4 when the report comes out, that one person would
5 say maybe I will get checked out. That can make a
6 difference. Thank you very much.

7 DR. MIDDENDORF: Thank you very much.

8 Next presenter is R.J. Lee.

9 R.J. LEE: I do want to thank the committee for
10 giving people the opportunity to testify. I've
11 been asked on behalf of the Policemen's Benevolence
12 Association to speak on their behalf about the
13 composition of the World Trade Center dust and some
14 analysis we recently did on the uniform of one
15 Officer Harris.

16 By way of background, R.J. Lee group worked in New
17 York City for about four years following the
18 disaster, characterizing, analyzing and
19 characterizing samples of World Trade Center dust
20 and exposures and things like that.

21 Today I want to talk about Officer Harris.

22 Laboratory testing of Officer Harris's clothing
23 worn on the morning of September 11th, clearly
24 demonstrates the presence of what's now referred to
25 as World Trade Center dust. And you can see the

1 uniform on the first slide that he was wearing that
2 day.

3 Fortunately, almost by, I don't know what fate,
4 Officer Harris had the presence of mind to go home
5 that morning and double bag his clothes so we have
6 a virgin sample of World Trade Center dust. One
7 that hadn't sat out in the rain, whatever, for
8 months, and one that you could look at as it was
9 created.

10 As you can see from what's called the World Trade
11 Center well, the World Trade Center dust is a
12 unique mixture of heavy metals, asbestos, fine
13 cement dust and chemicals produced by burning,
14 including PCBs, dioxins and furans. The chemical
15 species found in WTC, chemical and physical
16 species, found in World Trade Center dust can cause
17 many harmful effects on the body including effects
18 on the nervous system, kidneys and cancer.

19 It's, as you've heard it's widely believed that
20 there's been an insufficient amount of time to
21 assess the potential for increased cancer risk.
22 However, I believe there's certainly reason to
23 assume that the acute exposure experienced by first
24 responders are significant and unique.

25 There are a number of factors to be considered that

1 could play a role in increased cancer risk to
2 individuals and the potential for more rapid
3 progression than you would expect.

4 First of all, the initial dose, acute exposure was
5 enormous.

6 Next slide? This is the dust we found on Officer
7 Harris's clothing. You'll note that in something
8 like two or three hours, about 59,000 structures
9 per centimeter squared had been deposited on his
10 clothes. Chromium was at 347 micrograms per foot
11 square. That's a lot in a two or three-hour
12 exposure. If you put that cast an imaginary
13 membrane through the breathing zone, you can
14 translate that kind of deposition rate into
15 exposures and they're large.

16 There's an abundance of respirable particles in the
17 dust, far more than ordinary. What's interesting,
18 and one of the prior speakers mentioned it, in the
19 analysis we did of these hundred thousand samples,
20 and including Officer Harris, many of them were
21 coated. The asbestos was coated with lead; the
22 asbestos was coated with mercury. The machines
23 don't analyze for dioxins in the electron
24 microscope but obviously dioxins and PCBs were
25 there.

1 DR. MIDDENDORF: One minute, please.

2 R.J. LEE: The presence of dust on Officer Harris's
3 uniform clearly demonstrates that the first
4 responders were exposed to extreme conditions.
5 There was reason to believe that you could
6 postulate a model in which the dust carried, the
7 caustic cement dust, carried toxins and those
8 toxins and that interaction of the pH 11 or 12
9 cement dust could well interact with the lungs and
10 deliver toxins much more rapidly than believed
11 possible.

12 I think it's important on behalf of the PBA to say
13 that given the service of the first responders that
14 we've heard about today and the trauma they're
15 going through, that any potential disease that
16 could be covered should be covered on their behalf.
17 And secondly the information they're seeking from
18 the City and the government should be released
19 anonymously so that it can be used scientifically.
20 With that I thank you.

21 DR. MIDDENDORF: Our last commenter is Philip
22 Landrigan.

23 PHILIP LANDRIGAN: Good afternoon, Madam Chairman.
24 I'm Philip Landrigan, I'm a physician and
25 occupational doctor. Chairman of the Department of

1 Preventive Medicine, Dean for Global Health at Mt.
2 Sinai School of Medicine. For six years I directed
3 the Division of Surveillance Hazard Evaluations and
4 Field Study at NIOSH, so in other words for those
5 six years, 1979 to 1985, I directed the National
6 Occupational Epidemiology Program for the United
7 States of America. So we, we know for a certainty
8 from multiple lines of evidence, that you've heard
9 a great deal of data here today, and I thought that
10 testimony presented just now about the contaminated
11 police uniform was striking. We know that the
12 responders to 9/11 were exposed to a complex mix of
13 known and suspect human carcinogens. We know that
14 the air sampling data that were collected
15 undercount the true level of contamination. I
16 think the testimony just heard substantiates that,
17 but it stands to logic anyway that there were no
18 sampling units extant in the first hours and days
19 after the attack when the concentrations were
20 highest, so we know that the responders were,
21 especially those who were caught in the dust cloud,
22 were exposed to unprecedentedly high levels of
23 airborne contaminants.
24 Now, our group at the Mt. Sinai School of Medicine,
25 in partnership with people at UMBNJ, Stony Brook,

1 Queens College, North Shore LOIJ and Bellevue have
2 just completed an epidemiologic analysis based on
3 approximately 20,000 responders, and we looked
4 specifically at cancer in them. This is an
5 analysis that follows on our earlier studies
6 showing persistence of lung disease and mental
7 health problems and GERD in the responders.

8 I'm not going to present great detail because it's
9 going to be submitted for publication in the next
10 couple or three days, but I am going to give you a
11 broad sketch of the findings.

12 Overall we found approximately a 14-percent excess
13 in cancer at all sites combined in this population,
14 and we found statistically significant excesses of
15 thyroid, prostate and hematolymphatic,

16 hematolymphopoietic cancers, in this population.

17 In broad outline our findings parallel the findings
18 that were released on September 10th of this year,
19 that they would present from the fire department.

20 It's, I think, the 14-percent excess in overall
21 cancer is striking given that in this population,
22 we had a 58 prevalent -- 58-percent prevalence of
23 never smokers, and we had sharp deficits for lung
24 cancer and laryngeal cancer and yet despite those
25 deficits in some of the most common cancers, we had

1 an overall excess incidence of cancer in the
2 population. These are striking findings.
3 Going back to your taxonomy this morning of the
4 straw poll, I think we've reached a point where, to
5 use Steve Markowitz's phrase, we can say with a
6 high degree of certainty that the exposures that
7 the responders experienced down there at Ground
8 Zero, and at the other World Trade Center sites,
9 can be said to -- we can reasonably anticipate that
10 those exposures are going to cause cancer.

11 So I think, I think it puts you in a very difficult
12 policies (sic), but you clearly don't have the kind
13 of epidemiologic proof that you would like to have
14 to declare with 95 percent certainty that there's a
15 cause and effect relationship here. We're not
16 going to be there for some time yet. But you have
17 to bear in mind that in legal cases, you don't have
18 to get to 95 percent; you have to get to
19 51 percent. It has to be more likely than not that
20 the exposure caused the disease. And I think we're
21 at, or very close to that point.

22 And what I'd like to ask you as members of this
23 committee to weigh that as you make your decision.
24 Thank you.

25 DR. MIDDENDORF: Thank you very much,

1 Dr. Landrigan.

2 You have about 15 minutes left.

3 DR. DEMENT (via telephone): This is John Dement.
4 I'm going to have to leave the meeting so I just
5 want to make that note.

6 DR. MIDDENDORF: Okay. Thank you very much.

7 DR. WARD: So Virginia, are you still on the line?

8 DR. WEAVER (via telephone): Yes, I am.

9 DR. WARD: So I did want to give the committee an
10 opportunity if they had any questions or comments
11 on Virginia's presentation.

12 (no response)

13 DR. WARD: Okay, so --

14 DR. TALASKA: Oh, I have one question, if I may. I
15 have one question.

16 DR. WEAVER: Okay.

17 DR. TALASKA: You mentioned a statement early on
18 when you were talking about the VOCs, about that
19 when the levels became, quote, extremely high, that
20 people were removed from the area. And I just have
21 to ask was the concern -- you know if the concern
22 for that was because of explosion?

23 DR. WEAVER: I don't know.

24 DR. TALASKA: Didn't say it in the paper.

25 DR. WEAVER: I don't think so but I was reading

1 seriously in the last week and I could have missed
2 it, and perhaps others on the committee who spent
3 more time with these data could weigh in.

4 DR. TALASKA: Thank you, though.

5 DR. MARKOWITZ: So I have another question for
6 Virginia. So in your experience working with
7 firefighters from previous studies, how common is
8 it to find benzene at fires?

9 DR. WEAVER: It's extraordinarily common. We often
10 use data that's now rather old but still very valid
11 about the components, the VOCs in smoke; and in one
12 study conducted by Harvard, benzene was present in
13 about 92 percent of smoke samples obtained. And
14 it's routinely found at levels well above the OSHA
15 panel. Butadiene is also very common as a
16 combustion product.

17 DR. HARRISON: This is not really a question for
18 Virginia, just maybe an observation and a prelude
19 to further discussion that we'll have. I guess I
20 haven't heard anything from the presentations today
21 that would lead me to understand that there was a
22 minimum dose or duration of exposure that we could
23 identify from the knowledge that we have to draw a
24 line.

25 I think it gets, you know, back to maybe something

1 that, Liz, you presented earlier about latency and
2 duration of exposure. I guess I just would throw
3 that out there just for an observation, that we
4 really don't have, based on the limited amount of
5 exposure data, you know, that we have from the
6 site, the fact that it wasn't captured in the first
7 several days, a way to define a minimum length or
8 vocation related to the occurrence of cancer.

9 DR. WARD: So there is one question for
10 Dr. Landrigan.

11 DR. MIDDENDORF: Yes, well, there was one question.

12 DR. WARD: Is he still there? Dr. Landrigan?

13 Okay, so would someone like to ask a question of
14 Dr. Landrigan?

15 DR. TALASKA: Thanks for coming back, Phil.

16 DR. LANDRIGAN: No problem.

17 DR. TALASKA: I was wondering if you had done any
18 analysis on the subset of people who were on the
19 Pile early on relative to the whole group.

20 DR. LANDRIGAN: Yeah, we tried to do that. We
21 certainly, in our previous paper that you've
22 probably seen, the one that was published in
23 September in Lancet, we saw a very clear gradients
24 in most diseases according to intensity of
25 exposure.

1 The people who were caught in the cloud had the
2 highest rates of pretty much every disease we
3 looked at; the people who arrived in the first 48
4 hours but missed the cloud were the second highest,
5 and then on down through several more gradations.
6 We saw that for most types of lung disease, most
7 mental health problems, for GERD. It was not so
8 striking for cancer. And it may be because of
9 smaller numbers of cases. Thank you. That's it?
10 Yeah, thank you.

11 **DISCUSSION ON PRESENTATIONS**

12 DR. WARD: So, I guess we're close to the end of
13 our day. And I guess one, it was suggested earlier
14 that maybe we look separately at the question of
15 biologic plausibility and the likelihood of cancer
16 but I think one of the issues I'm struggling with,
17 and I don't know if other members of the committee
18 are struggling with it, too, is that we are --
19 whatever opinion we come to, we do have to define a
20 scientific rationale, and I know that in a lot of
21 the presentations this morning, you know, it would
22 be more possible to build a scientific rationale
23 around upper respiratory cancer, lung cancer,
24 esophageal cancer, areas of the body where we know
25 that there was direct contact with the carcinogenic

1 substances and we know that there have been other
2 kind of health effects, but I think the
3 difficulties we, we don't -- I mean, I guess, and
4 maybe Dr. Landrigan's study will help with that but
5 with the hematologic cancers and the lymphomas, we
6 don't as yet, I think, have strong epidemiologic
7 evidence, and I'm not sure we have, you know, an
8 exposure -- you know, we have a strong argument in
9 terms of biologic plausibility, and I guess -- so
10 the argument about -- I think we can say that, you
11 know, it's in shorter -- it's observed that they
12 have a shorter latency period but in terms of -- so
13 I guess what I'm seeking is, are that -- do people
14 have thoughts on that. How should we approach the
15 question of the blood cancers given that that seems
16 to be something that people are highly concerned
17 about? Excuse me? Does anyone care to comment on
18 that?

19 DR. WEAVER: So this is Virginia, and you know,
20 blood cancers are the ones that based on latency
21 alone, we could be seeing now from World Trade
22 Center exposures. You know, ten years out, those
23 would be the first wave of cancers that you would
24 see. Those are also caused, or closely connected,
25 with a number of the VOCs. And if you look at VOCs

1 in combustion products, they ask -- there are a
2 number. So you have an exposure mixture going on
3 there. And so from that point of view, I can see
4 the biological plausibility and that being an
5 initial concern.

6 DR. ROM: I think by definition, volatile means
7 volatile, that these compounds probably were very
8 high, right at the beginning with the burning of
9 all the fuels, and they evaporated into the air and
10 they weren't measured, and exposures were probably
11 way higher than any of the standards so that it's
12 biologically plausible that you're going to see
13 non-Hodgkin's, Hodgkin's lymphomas and the acute
14 leukemias, acute myelogenous or non-lymphatic
15 leukemia and probably chronic myelogenous leukemia.
16 I think the ALL and CLL are different biologies,
17 and that may be something totally different 'cause
18 ALL is in children and CLL is in the elderly
19 associated with a lot of genetic mutation defects.
20 But the others, and multiple myeloma, I would add,
21 probably all are very biologically plausible at
22 this time.

23 DR. MARKOWITZ: Also the firefighters study in fact
24 was positive for non-Hodgkin's lymphoma. It showed
25 a relative risk of 1.58 -- and actually whether you

1 use the corrected one, which tries to take account
2 of the surveillance issue or not, it showed a 50-
3 to 60-percent increase when compared to the general
4 population of men, and when they looked at it
5 compared to the firefighters who hadn't been
6 exposed, it was still elevated; it was 80- to 90-
7 percent increase. Not statistically significant at
8 that point because the numbers are smaller, but
9 when it was compared to the general population it
10 was elevated and that was statistically
11 significant, so there was real epidemiologic
12 evidence that blood cancer was increased.

13 DR. TALASKA: I think we might want to look more,
14 too, at some of the other compounds that we haven't
15 really spent any time with: the furans, the
16 dioxins; what sort of impact they have, both
17 animals and -- in animal studies for the most part,
18 to see if there is a link between those -- or
19 perhaps an interaction between those. And I don't
20 think anyone has looked at those as hard as maybe
21 we should.

22 DR. ALDRICH: (Indiscernible) the document that's
23 not biological plausibility (indiscernible).
24 Mesothelioma sometime in the distant future and
25 probably lung cancer in a little bit less distant

1 future, relative to the asbestos exposure. It's
2 hard to quantify but certainly potentially a
3 factor.

4 The fire department study did not show an increase
5 in lung cancer; it actually showed a decrease in
6 lung cancer possibly related to the health worker
7 effect, but that was seven years of study, and that
8 was probably too early to see the effects.

9 DR. WARD: So I guess I'm getting a sense. I know
10 some people have not spoken very much today but the
11 sense of the comments I'm getting is that many
12 people on the committee feel that it is certainly
13 biologically plausible that we would be seeing some
14 cancers in excess, either now or in the future, and
15 I guess the question is, is there someone who wants
16 to state, you know, make a statement -- or are
17 there people who would like to speak to the
18 question who have not spoken on it? Or we can go
19 back to the, you know, the poll, but I guess I'm
20 just trying to get a sense of the committee, of
21 where we stand at this point. Time, again, so we
22 can think about how we want to frame the discussion
23 tomorrow in the maximal -- you know, in a
24 productive way. Valerie?

25 MS. DABAS: Just from my observation, I understand

1 that the latency period for blood cancers is short.
2 I think we get into a very funny situation when we
3 start piecemealing each part out. Both the fire
4 study and Mt. Sinai seem to indicate that thyroid
5 and prostate, they're seeing increases, and so if
6 we start going by what is easiest and not looking
7 at the whole picture, then I think we may start
8 asking too -- well, I guess you can't ask too many
9 questions but then it gets very confusing.

10 For me, I've seen, you know, from taking
11 information from responders, I've seen an increase
12 in thyroid, I've seen an increase in prostate. I
13 was told that, you know, thyroid is common,
14 prostate is common, but when we look at the ages
15 people are being diagnosed, it's very uncommon for
16 a 38-year-old man to even be tested for prostate
17 cancer, so when they come up with prostate cancer,
18 I think it's significant.

19 I also have seen an increase -- you know, how do
20 you deal, then, with the blood and liver canc --
21 kidney cancers that we're seeing? Liver cancers
22 with people that are not hepatitis C and do not
23 have cirrhosis of the liver. You know, we had four
24 cases reported in that instance and, you know, so
25 you have to really look at the whole picture as

1 opposed to just saying well, the blood cancers are
2 a four-to-six year latency period, we're at four to
3 six years. If that's the case, that's just
4 assuming that the dust is the same exposure as
5 we've seen with all these other studies, and I
6 don't think these studies take into effect the
7 concentration of chemicals, metals and so forth,
8 and we keep saying the dust is different than
9 anything that we've seen before, and therefore I
10 think we have to treat it different.

11 MR. CASSIDY: I just wanted to add that I think
12 it's clear that we need to remember what was
13 highlighted today, which is that this type of
14 exposure to the variety of different things, the
15 concrete, the dust, the metals, the benzene, all
16 the chemicals, really hasn't been -- we haven't
17 seen that anywhere before so when you want to start
18 breaking down studies and say well, exposure to
19 benzene means this. When you add them all
20 together, you really have a toxic stew that, I
21 think, is so biologically plausible to say that
22 blood cancers and these other cancers are a result
23 of that exposure, and I do think the severity of
24 the exposure, you know, bears out clearly that, you
25 know, those who were caught in the dust, in the

1 cloud, in the collapse, those who were there in the
2 48 hours, those who spent extensive times there,
3 clearly have a more likely coming down with these
4 cancers, but I think it's biologically plausible
5 that anyone that was subject to this is going to
6 have an increased rate of cancer so that my view
7 now, given everything that I've heard, is that that
8 cancer should be included.

9 We need a better mic system.

10 DR. HARRISON: Steve, this is Bob Harrison. Were
11 you saying that we should recommend that all
12 cancers be covered regardless of site?

13 MR. CASSIDY: I'm sorry? I think to say all is a
14 broad statement; it really is. But I think that
15 clearly the blood cancers, which are showing up
16 early, I think anything related to the lungs, the
17 respiratory system, anything that you can possibly
18 inhale, so the esophageal cancers. You know, the
19 fire department study proves that firefighters lost
20 12 years' lung capacity in the blink of an eye.
21 That can't be dismissed as -- if that didn't exist
22 people would say well, maybe this dust cloud really
23 isn't going to do anything to us. But it proved
24 what happened. Twelve years lung capacity, so to
25 say all? I'm not saying all but I think we should

1 err on the side of, if there's any evidence, we
2 should err on that side.

3 MS. FLYNN: I really appreciated Dr. Landrigan's
4 comments, and I just want to say that I think that
5 this is obviously not a deliberation that should
6 use, you know, scientific certainty; this has been
7 said before.

8 As his basis, he talked about a 51-percent of, you
9 know, using the phrase that Steve Markowitz used
10 earlier: We can reasonably anticipate that these
11 cancers are linked to World Trade Center exposures,
12 and right now that sounds pretty right to me.

13 I also want to add that the community cannot be
14 left out of this deliberation, and also that the
15 James Zadroga Act, and I can provide pages to folks
16 if they want them, provides for one list of World
17 Trade Center-covered conditions.

18 And we all know as erratic and full of gaps as the
19 sampling information was on the Pile, you know, how
20 much more is not known about community exposures.

21 But what we do know is that members of the
22 community, residents, students and area workers
23 have the same respiratory and the same set of
24 aerodigestive 9/11-related illnesses as responders,
25 and it's more than reasonable to anticipate that

1 they would develop the same set of cancers.

2 MS. HUGHES: I also just wanted to -- I'm not a
3 biology expert, but I did go online and if we could
4 break the body down into different body systems,
5 like respiratory, and then look at the different
6 things that could be impacted, so it is not just
7 necessarily the lungs but it's the throat, so we're
8 looking at a comprehensively wide body system so I
9 just wanted to add that as well.

10 **ADMINISTRATIVE ISSUES AND ADJOURN**

11 DR. WARD: So we do need to leave the building
12 shortly. So again I'm trying to sum up the sense
13 that I'm getting. It seems that many people are in
14 favor of listing at least some cancers of some
15 systems as World Trade Center-related conditions,
16 so I guess, you know, your homework assignment is
17 to really maybe clarify your own position as much
18 as possible, and try to come up with potential
19 statements that you think the group could agree on,
20 and y'all certainly be thinking about it, but I'd
21 like, you know, others as well to come in with, I
22 think this is the sense of the committee and we can
23 capture it in these words. That would really I
24 think move us along in the morning.
25 So well, I did want to thank everyone who's here,

1 both those who spoke and those who did not speak.
2 I think, you know, the public comments are very
3 informative. I think the discussion today was very
4 informative, and I hope we've moved towards --
5 we've moved forward in the process of making a
6 recommendation.

7 DR. MIDDENDORF: Let me also express my thanks and
8 thanks for NIOSH and the World Trade Center Health
9 Program, for the participation of everyone.

10 Steve, your wish is our command. We will be in
11 conference rooms A and B tomorrow. And the speaker
12 system will be better. It's not perfect but it
13 will be better. So for any members of the public
14 who intend to come back, we will be at the other
15 end on the same floor. Thank you and good night.

16 (Meeting adjourned for the day at 5:05 p.m.)
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CERTIFICATE OF COURT REPORTER
STATE OF GEORGIA
COUNTY OF FULTON

I, Steven Ray Green, Certified Merit Master Court Reporter, do hereby certify that I reported the above and foregoing on the day of February 15, 2012; and it is a true and accurate transcript of the proceedings captioned herein.

I further certify that I am neither related to nor counsel to any of the parties herein, nor have any interest in the cause named herein.

WITNESS my hand and official seal this the 9th day of March, 2012.

STEVEN RAY GREEN, CCR, CVR-CM-M, PNSC
CERTIFIED MERIT MASTER COURT REPORTER
CERTIFICATE NUMBER: A-2102

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